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(54) Title: INHIBITORS OF INTERLEUKIN-Iβ CONVERTING ENZYME

57) Abstra

The present invention relates to novel classes of compounds which are inhibitors of interleukin-19 converting enzyme. The LOE inhibitors of this invention are characterized by specific structural and physicochemical features. This invention also relates to make the compositions comprising these compounds. The compounds and pharmaceutical compositions of this invention are particularly well suited for inhibiting ICE activity and consequently, may be advantageously used as agents against III-1 appoints[5]. IGIP—and interest diseases, autoimmune diseases, destructive bone disorders, proliferative disorders, infectious diseases, degenerative diseases, and energies diseases, and energies of the control of the compounds of the compounds of the compounds of the compounds and compositions of this invention. This invention also relates to methods for inhibiting ICE activity, for treating interclavit, apoptosis, profits invention also relates to methods for inhibiting also relates to methods for inhibiting also relates to methods for inhibiting also relates to methods for preparing N-acytamino compounds.

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INHIBITORS OF INTERLEUKIN-18 CONVERTING ENZYME

TECHNICAL FIELD OF THE INVENTION

The present invention relates to novel 5 classes of compounds which are inhibitors of interleukin-18 converting enzyme ("ICE"). This invention also relates to pharmaceutical compositions comprising these compounds. The compounds and pharmaceutical compositions of this invention are 10 particularly well suited for inhibiting ICE activity and consequently, may be advantageously used as agents against interleukin-1- ("IL-1"), apoptosis-, interferon gamma inducing factor- ("IGIF") and interferon-v-("IFN-y") mediated diseases, including inflammatory 15 diseases, autoimmune diseases, destructive bone. proliferative disorders, infectious diseases and degenerative diseases. This invention also relates to methods for inhibiting ICE activity, and decreasing IGIF production and IFN-y production and methods for 20 treating interleukin-1-, apoptosis-, IGIF- and IFN-ymediated diseases using the compounds and compositions of this invention. This invention also relates to

methods of preparing N-acylamino compounds.

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BACKGROUND OF THE INVENTION

Interleukin 1 ("IL-1") is a major proinflammatory and immunoregulatory protein that stimulates fibroblast differentiation and proliferation, the production of prostaglandins, 5 collagenase and phospholipase by synovial cells and chondrocytes, basophil and eosinophil degranulation and neutrophil activation. Oppenheim, J.H. et al, Immunclogy Today, 7, pp. 45-56 (1986). As such, it is involved in the pathogenesis of chronic and acute 1.0 inflammatory and autoimmune diseases. For example, in rheumatoid arthritis, IL-1 is both a mediator of inflammatory symptoms and of the destruction of the cartilage proteoglycan in afflicted joints. Wood, D.D. et al., Arthritis Rheum. 26, 975, (1983); Pettipher, 15 E.J. et al., Proc. Natl. Acad. Sci. UNITED STATES OF AMERICA 71, 295 (1986); Arend, W.P. and Dayer, J.M., Arthritis Rheum. 38, 151 (1995). IL-1 is also a highly potent bone resorption agent. Jandiski, J.J., J. Oral Path 17, 145 (1988); Dewhirst, F.E. et al., J. Immunol. 20 8, 2562 1985). It is alternately referred to as "osteoclast activating factor" in destructive bone diseases such as osteoarthritis and multiple myeloma. Bataille, R. et al., Int. J. Clin. Lab. Res. 21(4), 283 25 (1992). In certain proliferative disorders, such as acute myelogenous leukemia and multiple myeloma, IL-1 can promote tumor cell growth and adhesion. Bani, M.R., J. Natl. Cancer Inst. 83, 123 (1991); Vidal-Vanaclocha, F., Cancer Res. 54, 2667 (1994). In these disorders. IL-1 also stimulates production of other 30 cytokines such as IL-6, which can modulate tumor development (Tartour et al., Cancer Res. 54, 6243 (1994). IL-1 is predominantly produced by peripheral blood monocytes as part of the inflammatory response

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and exists in two distinct agonist forms, IL-1 α and IL-1 β . Mosely, B.S. et al., <u>Proc. Nat. Acad. Sci.</u>, 84, pp. 4572-4576 (1987); Lonnemann, G. et al., <u>Eur.J. Immunol.</u>, 19, pp. 1531-1536 (1989).

IL-1β is synthesized as a biologically inactive precursor, pIL-1β. pIL-1β lacks a conventional leader sequence and is not processed by a signal peptidase. March, C.J., Nature, 315, pp. 641-647 (1985). Instead, pIL-1β is cleaved by interleukin-1β converting enzyme ("ICE") between Asp-116 and Ala-117 to produce the biologically active C-terminal fragment found in human serum and synovial fluid. Sleath, P.R., et al., J. Biol. Chem., 265, pp. 14526-14528 (1992); A.D. Howard et al., J.

155 Immunol., 147, pp. 2964-2969 (1991). ICE is a cysteine protease localized primarily in monocytes. It converts precursor IL-1β to the mature form. Black, R.A. et al., FEBS Lett., 247, pp. 386-390 (1989); Kostura, M.J. et al., Proc. Natl. Acad. Sci. UNITED STATES OF AMERICA, 86, pp. 5227-5231 (1989). Processing by ICE is also necessary for the transport of mature IL-1β

through the cell membrane.

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ICE, or its homologs, also appears to be involved in the regulation of programmed cell death or apoptosis. Yuan, J. et al., Cell, 75, pp. 641-652 (1993); Miura, M. et al., Cell, 75, pp. 653-660 (1993); Nett-Fiordalis:, M.A. et al., J. Cell Biochem., 17B, p. 117 (1993). In particular, ICE or ICE homologs are thought to be associated with the regulation of

apoptosis in neurodegenerative diseases, such as Alzheimer's and Parkinson's disease. Marx, J. and M. Baringa, <u>Science</u>, 259, pp. 760-762 (1993); Gagliardini, V. et al., <u>Science</u>, 263, pp. 826-828 (1994). Therapeutic applications for inhibition of apoptosis

may include treatment of Alzheimer's disease,
Parkinson's disease, stroke, myocardial infarction,
spinal atrophy, and aging.

ICE has been demonstrated to mediate

apoptosis (programmed cell death) in certain tissue
types. Steller, H., Science, 267, p. 1445 (1995);
Whyte, M. and Bvan, G., Nature, 376, p. 17 (1995);
Martin, S.J. and Green, D.R., Cell, 82, p. 349 (1995);
Alnemri, E.S., et al., J. Biol. Chem., 270, p. 4312
(1995); Yuan, J. Curr. Opin. Cell Biol., 7, p. 211
(1995). A transgenic mouse with a disruption of the
ICE gene is deficient in Fas-mediated apoptosis (Kuida,

ICE is distinct from its role as the processing enzyme for pro-IL1\(\beta\). It is conceivable that in certain tissue types, inhibition of ICE may not affect secretion of mature IL-1\(\beta\), but may inhibit apoptosis.

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K. et al., Science 267, 2000 (1995)). This activity of

Enzymatically active ICE has been previously described as a heterodimer composed of two subunits, p20 and p10 (20kDa and 10kDa molecular weight, respectively). These subunits are derived from a 45kDa proenzyme (p45) by way of a p30 form, through an activation mechanism that is autocatalytic. Thornberry, N.A. et al., Nature, 356, pp. 768-774 (1992). The ICE proenzyme has been divided into several functional domains: a prodomain (p14), a p22/20 subunit, a polypeptide linker and a p10 subunit. Thornberry et al., supra; Casano et al., Genomics, 20, pp. 474-481 (1994).

Full length p45 has been characterized by its cDNA and amino acid sequences. PCT patent applications WO 91/15577 and WO 94/00154. The p20 and p10 cDNA and amino acid sequences are also known. Thornberry et al., supra. Murine and rat ICE have also been sequenced and cloned. They have high amino acid and

nucleic acid sequence homology to human ICE. Miller, D.K. et al., Ann. N.Y. Acad. Sci., 696, pp. 133-148 (1993); Molineaux, S.M. et al., Proc. Nat. Acad. Sci., 90, pp. 1809-1813 (1993). The three-dimensional structure of ICE has been determined at atomic resolution by X-ray crystallography. Wilson, K.P., et al., Nature, 370, pp. 270-275 (1994). The active enzyme exists as a tetramer of two p20 and two p10

Additionally, there exist human homologs of ICE with sequence similarities in the active site regions of the enzymes. Such homologs include TX (or ICE_{rel-II} or ICH-2) (Faucheu, et al., <u>EMBO J.</u>, 14, p. 1914 (1995); Kamens J., et al., <u>J. Biol. Chem.</u>, 270, p. 15 15250 (1995); Nicholson et al., <u>J. Biol. Chem.</u>, 270

subunits.

20 (1995), CPP32 (or YAMA or apopain) (Fernandes-Alnemri, T. et al., J. Biol. Chem., 269, p. 30761 (1994); Nicholson, D.W. et al., Nature, 376, p. 37 (1995)), and CMH-1 (or MCH-3) (Lippke, et al., J. Biol. Chem., (1996); Fernandes-Alnemri, T. et al., Cancer Res.,

25 (1995)). Each of these ICE homologs, as well as ICE itself, is capable of inducing apoptosis when overexpressed in transfected cell lines. Inhibition of one or more of these homologs with the peptidyl ICE inhibitor Tyr-Val-Ala-Asp-chloromethylketone results in

inhibition of apoptosis in primary cells or cell lines.

Lazebnik et al., Nature, 371, p. 346 (1994). The compounds described herein are also capable of inhibiting one or more homologs of ICE (see Example 5). Therefore, these compounds may be used to inhibit

35 apoptosis in tissue types that contain ICE homologs, but which do not contain active ICE or produce mature

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IL-1B.

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Interferon-gamma inducing factor (IGIF) is an approximately 18-kDa polypeptide that stimulates T-cell production of interferon-gamma (IFN-y). IGIF is produced by activated Kupffer cells and macrophages in vivo and is exported out of such cells upon endotoxin stimulation. Thus, a compound that decreases IGIF production would be useful as an inhibitor of such T-cell stimulation which in turn would reduce the levels of IFN-y production by those cells.

IFN-y is a cytokine with immunomodulatory effects on a variety of immune cells. In particular, IFN-y is involved in macrophage activation and Th1 cell selection (F. Belardelli, AFMIS, 103, p. 161 (1995)). IFN-y exerts its effects in part by modulating the expression of genes through the STAT and IFF pathways (C. Schindler and J.E. Darnell, Ann. Rev. Biochem., 64, p. 621 (1995); T. Taniguchi, J. Cancer Res. Clin. Oncol., 121, p. 516 (1995)).

Mice lacking IFN-y or its receptor have multiple defects in immune cell function and are resistant to endotoxic shock (S. Huang et al., <u>Science</u>, 259, p. 1742 (1993); D. Dalton et al., <u>Science</u>, 259, p. 1739 (1993); B. D. Car et al., <u>J. Exp. Med.</u>, 179, p. 1437 (1994)). Along with IL-12, IGIF appears to be a potent inducer of IFN-y production by T cells (H. Okamura et al., <u>Infection and Immunity</u>, 63, p. 3966 (1995); H. Okamura et al., <u>Nature</u>, 378, p. 88 (1995); S. Ushio et al., <u>J. Immunol.</u>, 156, p. 4274 (1996)).

IFN-y has been shown to contribute to the pathology associated with a variety of inflammatory, infectious and autoimmune disorders and diseases.

Thus, compounds capable of decreasing IFN-y production

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would be useful to amelicrate the effects of IFN- $_{\rm Y}$ related pathologies.

The biological regulation of IGIF and thus IFN-Y has not been elucidated. It is known that IGIF is synthesized as a precursor protein, called "pro-IGIF". It has been unclear, however, how pro-IGIF is cleaved and whether its processing has biological importance.

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Accordingly, compositions and methods capable
of regulating the conversion of pro-IGIF to IGIF would
be useful for decreasing IGIF and IFN-y production in

<u>Vivo</u>, and thus for ameliorating the detrimental effects
of these proteins which contribute to human disorders
and diseases.

15 However, ICE and other members of the ICE/CED-3 family have not previously been linked to the conversion of pro-IGIF to IGIF or to IFN-γ production in vivo.

ICE inhibitors represent a class of compounds 20 useful for the control of inflammation or apoptosis or both. Peptide and peptidyl inhibitors of ICE have been described. PCT patent applications WO 91/15577; WO 93/05071; WO 93/09135; WO 93/14777 and WO 93/16710; and European patent application 0 547 699. Such peptidyl 25 inhibitors of ICE has been observed to block the production of mature $IL-1\beta$ in a mouse model of inflammation (vide infra) and to suppress growth of leukemia cells in vitro (Estrov et al., Blood 84, 380a (1994)). However, due to their peptidic nature, such inhibitors are typically characterized by undesirable 3.0 pharmacologic properties, such as poor cellular penetration and cellular activity, poor oral absorption, poor stability and rapid metabolism. Plattner, J.J. and D.W. Norbeck, in Drug Discovery

- 8 -

<u>Technologies</u>, C.R. Clark and W.H. Moos, Eds. (Ellis Horwood, Chichester, England, 1990), pp. 92-126. This has hampered their development into effective drugs.

Non-peptidyl compounds have also been reported to inhibit ICE in vitro. PCT patent application WO 95/26958; US Patents 5,552,400; Dolle et al., J. Med. Chem., 39, pp. 2438-2440 (1996); However, it is not clear whether these compounds have

the appropriate pharmacological profile to be therapeutically useful.

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Additionally, current methods for the preparation of such compounds are not advantageous. These methods use tributyltin hydride, a toxic, moisture sensitive reagent. Thus, these methods are inconvenient to carry out, pose a health risk and

15 inconvenient to carry out, pose a health risk and create toxic-waste disposal problems. Furthermore, it is difficult to purify compounds prepared by these methods.

Accordingly, the need exists for compounds

that can effectively inhibit the action of ICE in vivo,
for use as agents for preventing and treating chronic
and acute forms of IL-1-mediated diseases, apoptosis-,
IGIF-, or IFN-y-mediated diseases, as well as
inflammatory, autoimmune, destructive bone,

25 proliferative, infectious, or degenerative diseases. The need also exists for methods of preparing such compounds.

SUMMARY OF THE INVENTION

The present invention provides novel classes of compounds, and pharmaceutically acceptable derivatives thereof, that are useful as inhibitors of ICE. These compounds can be used alone or in combination with other therapeutic or prophylactic agents, such as antibiotics, immunomodulators or other anti-inflammatory agents, for the treatment or

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prophylaxis of diseases mediated by IL-1, apoptosis, IGIF or IFN-Y. According to a preferred embodiment, the compounds of this invention are capable of binding to the active site of ICE and inhibiting the activity of that enzyme. Additionally, they have improved cellular potency, improved pharmacokinetics, and/or improved oral bioavailability compared to peptidyl ICE inhibitors.

It is a principal object of this invention to provide novel classes of compounds which are inhibitors of ICE represented by formulas:

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(II)-(V) and (VII)
$$\bigcap_{m \text{ } OR_{13}} OR_{13}$$

wherein the various substituents are described herein.

It is a further object of this invention to provide a process of preparing N-acylamino compounds by coupling a carboxylic acid with an alloc-protected amine.

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BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1A ICE cleaves pro-IGIF in vivo. Cell lysates from Cos cells transfected with the various indicated expression plasmids or controls were analyzed for the presence of IGIF by separating proteins by SDS-PAGE and immunoblotting with anti-IGIF antisera (lane 1, mock transfected cells; lane 2, pro-IGIF alone; lanes 3-12, pro-IGIF in combination with ICE, ICE-C285S, CPP32, CPP32-C163S, CMH-1, CMH-1-C186S, Tx, Tx-C258S, respectively). Mobilities of pro-IGIF and the 18-kDa mature IGIF are indicated on the right. Molecular weight markers in kDa are shown on the left (Example 23).

Fig. 1B ICE cleaves pro-IGIF at the authentic 1.5 processing site in vitro as shown by Coomassie blue staining of proteolytic reaction products separated by SDS-PAGE (Example 23). The proteases and inhibitors used were: lane 1, buffer control; lane 2, 0.1 nM ICE: lane 3, 1 nM ICE; lanes 4 and 5, 1 nM ICE with 10 nM 20 Cbz-Val-Ala-Asp-[(2,6-dichlorobenzoyl)oxy]methyl ketone and 100 nM Ac-Tyr-Val-Ala-Asp-aldehyde, respectively; lanes 6 and 7, 15 nM CPP32 with and without 400 nM Ac-Asp-Glu-Val-Asp-aldehyde (D. W. Nicholson et al., Nature, 376, p. 37 (1995)), respectively; lane 8, 100 25 nM CMH-1; lane 9, 10 units/ml granzyme B; and M, molecular weight markers in kDa.

Fig. 1C ICE cleavage converts inactive pro-IGIF to active IGIF which induces IFN-y production in Th1 helper cells. Uncleaved (Pro-IGIF), ICE-cleaved (Pro-IGIF/CP32), and recombinant mature IGIF (rIGIF) were incubated with

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A.E7 Thi cells at 12 ng/ml (open bar) and 120 ng/ml (hatched bar) for eighteen hours and the levels of IFN-Y released into the culture medium assayed by ELISA (Example 23). A.E7 cells cultured with buffer, ICE alone (ICE) or CPP32 alone (CPP32) were assayed similarly for negative controls. The numbers represent the average of three determinations.

Fig. 2A Mature IGIF (18-kDa) is produced by Cos cells co-transfected with pro-IGIF and ICE-expressing plasmids. Cell lysates (left) and conditioned medium 1.0 (right) from Cos cells transfected with a pro-IGIF expression plasmid in the absence (-) or presence of an expression plasmid encoding wild type (ICE) or inactive mutant (ICE-C285S) ICE. Transfected cells were metabolically labeled with 35S-methionine, proteins from 15 cell lysates and conditioned medium immunoprecipitated with anti-IGIF antisera and separated by SDS-PAGE (Example 24). Mobilities of pro-IGIF and the 18-kDa mature IGIF are indicated on the right. Molecular 20 weight markers in kDa are shown on the left.

Fig. 2B IFN-v inducing activity is detected in Coscells co-transfected with pro-IGIF and ICE-expressing plasmids. Cell lysates (hatched bar) and conditioned medium (open bar) from Coscells transfected with a pro-IGIF expression plasmid in the absence (Pro-IGIF) or presence (Pro-IGIF/ICE) of an expression plasmid encoding wild type (ICE) were assayed for IFN-v levels (ng/ml) by ELISA. Coscells transfected with buffer (Mock) or an ICE-expressing plasmid alone (ICE) served as negative controls (Example 24).

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- Fig. 3A Kupffer cells from mice lacking ICE are defective in the export of IGIF. Kupffer cells from wild type mice (ICE +/+) or ICE-deficient mice homozygous for an ICE mutation (ICE-/-) were isolated and primed with LPS for three hours. The levels of immunoreactive IGIF polypeptides in the conditioned media (ng/ml) of wild type cells were measured by ELISA (Example 25). N.D. (not detectable) indicates that the IGIF concentration was less than 0.1 ng/ml.
- 10 Fig. 3B Kupffer cells from mice lacking ICE are defective in the export of mature IGIF. Kupffer cells from wild type mice (ICE +/+) or ICE deficient mice homozygous for an ICE mutation (ICE -/-) were isolated and primed with LPS for three hours. Primed cells were metabolically labeled with 35S-methionine, proteins from cell lysates and conditioned medium immunoprecipitated with anti-IGIF antisera and separated by SDS-PAGE (Example 25). Mobilities of pro-IGIF and the 18-kDa mature IGIF are indicated on the right. Molecular mass
 - Fig. 3C Serum from ICE-deficient mice contains reduced levels of IGIF. Serum samples from wild type mice (ICE +/+) or ICE deficient mice homozygous for an ICE mutation (ICE -/-) were assayed for IGIF levels (hg/ml) by ELISA (Example 25).
 - Fig. 3D Serum from ICE-deficient mice contains reduced levels of IFN-v. Serum samples from wild type mice (ICE +/+) or ICE deficient mice homozygous for an ICE mutation (ICE -/-) were assayed for IFN-v levels (ng/ml) by ELISA (Example 25).

- Fig. 4 Serum IFN-y levels are significantly reduced in ICE-deficient mice after an acute challenge with LPS (Example 26). Serum samples from wild type mice (filled squares) or ICE-deficient mice (filled circles) were assayed for IFN-y levels (ng/ml) by ELISA as a function of time (hours) after LPS challenge.

 Temperatures of the animals during the time course in degrees Celcius is shown for wild type mice (open squares) or ICE-deficient mice (open circles).
- 10 Fig. 5 The ICE inhibitor, AcYVAD-aldehyde (AcYVAD-CHO), inhibits LPS-stimulated IL-18 and IFN-y synthesis by human peripheral blood mononuclear cells (PBMC).

 Percent (%) inhibition as a function of inhibitor concentration (µM) is shown for IL-18 (open squares)

 15 and IFN-y (open diamonds) synthesis.
 - Fig. 6 Compound 214e inhibits IL-1β production in LPS-challenged mice. Serum samples from CD1 mice were assayed for IL-1β levels (pg/ml) by ELISA after LPS challenge. Compound 214e was administered by intraperitoneal (IP) injection one hour after LPS challenge. Blood was collected seven hours after LPS challenge (see Example 7).
- Fig. 7 Compound 217e inhibits IL-1β production in LPS-challenged mice. Serum samples from CD1 mice were assayed for IL-1β levels (pg/ml) by ELISA after LPS challenge. Compound 217e was administered by intraperitoneal (IP) injection one hour after LPS challenge. Blood was collected seven hours after LPS challenge (see Example 7).
- 30 Fig. 8 Compound 214e, but not compound 217e,

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inhibits IL-1 β production in LPS-challenged mice when administered by oral gavage. This assay measures oral absorption under similar conditions as those described for Figs. 6 and 7. These results indicates that 214e is potentially orally active as an ICE inhibitor (see Example 7).

- Fig. 9 Compound 214e and analogs of 214e also inhibit IL-1\$ production after IP administration. These results were obtained in the assay described for Figs. 6 and 7 and Example 7.
- Fig. 10 Compound 214e, and analogs of 214e, also inhibit IL-18 production after oral (PO) administration. These results were obtained in the assay described for Figs. 6 and 7 and Example 7.
- 15 Figs. 11A/B Compounds 302 and 304a show detectable blood levels when administered orally (50mg/kg, in 0.5 % carboxymethylcellulose) to mice. Blood samples were collected at 1 and 7 hours after dosing. Compounds 302 and 304a are prodrugs of 214e and are metabolized to 20 214e in vivo. Compound 214e shows no blood levels above 0.10 μg/ml when administered orally (Example 8).
- Fig. 12 Compound 412f blocks the progression of type II collagen-induced arthritis in male DBA/1J mice (Wooley, P.H., Methods in Enzymology, 162, pp. 361-373 (1988) and Geiger, T., Clinical and Experimental Rheumatology, 11, pp. 515-522 (1993)). Compound 412f was administered twice a day (10, 25 and 50mg/kg), approximately 7h apart, by oral gavage. Inflammation was measured on the Arthritis Severity Score on a 1 to 4 scale of increasing severity. The scores of the two front paws were added to give the final score (see Example 21).

- Fig. 13 Compound 412d blocks the progression of type II collagen-induced arthritis in male DBA/1J mice. The results were obtained as described for Fig. 12 and in Example 21.
- Fig. 14 Compound 696a blocks the progression of type II collagen-induced arthritis in male DBA/1J mice. The results were obtained as described for Fig. 12 and in Example 21.

ABBREVIATIONS AND DEFINITIONS

10	Abbreviations		
	Designation	Reagent or Fragment	
	Ala	alanine	
	Arg	arginine	
	Asn	asparagine	
15	Asp	aspartic acid	
	Cys	cysteine	
	Gln	glutamine	
	Glu	glutamic acid	
	Gly	glycine	
20	His	histidine	
	Ile	isoleucine	
	Leu	leucine	
	Lys	lysine	
	Met	methionine	
25	Phe	phenylalanine	
	Pro	proline	
	Ser	serine	
	Thr	threonine	
	Trp	tryptophan	
30	Tyr	tyrosine	
	Val	valine	
	Ac ₂ O	acetic anhydride	
	n-Bu	normal-butyl	

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	DMF	dimethylformamide
	DIEA	N, N-diisopropylethylamine
	EDC	1-(3-Dimethylaminopropyl)-3-
		ethylcarbodiimide hydrochloride
5	Et ₂ O	diethyl ether
	EtOAc	ethyl acetate
	Fmoc	9-fluorenylmethyoxycarbonyl
	HBTU	O-benzotriazol-1-yl-N, N, N', N'-
		tetramethyluronium
10		hexafluorophosphate
	HOBT	1-hydroxybenzotriazole hydrate
	MeOH	methanol
	TFA	trifluoroacetic acid
	Alloc	allyloxycarbonyl
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Definitions

The following terms are employed herein:
The term "interferon gamma inducing factor"
or "IGIF" refers to a factor which is capable of
stimulating the endogenous production of IFN-y.

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The term "ICE inhibitor" refers to a compound which is capable of inhibiting the ICE enzyme. ICE inhibition may be determined using the methods described and incorporated by reference herein. The skilled practitioner realizes that an <u>in vivo</u> ICE inhibitor is not necessarily an <u>in vitro</u> ICE inhibitor. For example, a prodrug form of a compound typically demonstrates little or no activity in <u>in vitro</u> assays. Such prodrug forms may be altered by metabolic or other biochemical processes in the patient to provide an <u>in vivo</u> ICE inhibitor.

 $\label{thm:cytokine} The \ term \ "cytokine" \ refers \ to \ a \ molecule \\ which \ mediates \ interactions \ between \ cells.$

The term "condition" refers to any disease,

disorder or effect that produces deleterious biological consequences in a subject.

The term "subject" refers to an animal, or to one or more cells derived from an animal. Preferably, the animal is a mammal, most preferably a human. Cells may be in any form, including but not limited to cells retained in tissue, cell clusters, immortalized cells, transfected or transformed cells, and cells derived from an animal that have been physically or

10 phenotypically altered.

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The term "active site" refers to any or all of the following sites in ICE: the substrate binding site, the site where an inhibitor binds and the site where the cleavage of substrate occurs.

The term "heterocycle" or "heterocyclic" refers to a stable mono- or polycyclic compound which may optionally contain one or two double bonds or may optionally contain one or more aromatic rings. Each heterocycle consists of carbon atoms and from one to four heteroatoms independently selected from a group including nitrogen, oxygen, and sulfur. As used herein, the terms "nitrogen heteroatoms" and "sulphur heteroatoms" include any oxidized form of nitrogen or sulfur and the quaternized form of any basic nitrogen. Heterocycles defined above include, for example, pyrimidinyl, tetrahydroquinolyl, tetrahydroisoquinonlinyl, purinyl, pyrimidyl,

tetrahydroisoquinonlinyl, purinyl, pyrimidyl, indolinyl, benzimidazolyl, imidazolyl, imidazolyl, imidazolinoyl, imidazolidinyl, quinolyl, isoquinolyl, indolyl, pyridyl, pyrrolyl, pyrrolyl, pyrazolyl, pyrazinyl, quinoxolyl, piperidinyl, morpholinyl, thiamorpholinyl, furyl, thienyl, triazolyl, thiazolyl, β-carbolinyl,

tetrazolyl, thiazolidinyl, benzofuranoyl, thiamorpholinyl sulfone, benzoxazolyl, oxopiperidinyl, oxopyrroldinyl, oxoazepinyl, azepinyl, isoxazolyl,

- 18 -

tetrahydropyranyl, tetrahydrofuranyl, thiadiazolyl, benzodioxolyl, benzothienyl, tetrahydrothiophenyl and sulfolanyl. Further heterocycles are described in A.R. Katritzky and C.W. Rees, eds., Comprehensive

Heterocyclic Chemistry: The Structure, Reactions, Synthesis and Use of Heterocyclic Compounds, Vol. 1-8, Pergamon Press, NY (1984).

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The term "cycloalkyl" refers to a mono- or polycyclic group which contains 3 to 15 carbons and may optionally contain one or two double bonds. Examples include cyclohexyl, adamantyl and norbornyl.

The term "aryl" refers to a mono- or polycyclic group which contains 6, 10, 12, or 14 carbons in which at least one ring is aromatic. Examples include phenyl, naphthyl, and tetrahydronaphthalene.

The term "heteroaromatic" refers to a monoor polycyclic group which contains 1 to 15 carbon atoms and from 1 to 4 heteroatoms, each of which is selected independently from a group including sulphur, nitrogen and oxygen, and which additionally contains from 1 to 3 five or six membered rings, at least one of which is aromatic.

The term "alpha-amino acid" (α -amino acid) refers to both the naturally occurring amino acids and other "non-protein" α -amino acids commonly utilized by those in the peptide chemistry arts when preparing synthetic analogues of naturally occurring peptides, including D and L forms. The naturally occurring amino acids are glycine, alanine, valine, leucine, isoleucine, serine, methionine, threonine, phenylalanine, tyrosine, tryptophan, cysteine, proline, histidine, aspartic acid, asparagine, glutamic acid, glutamine, y-carboxyglutamic acid, arginine, ornithine and lysine. Examples of "non-protein" alpha-amino acids include

hydroxylysine, homoserine, homotyrosine, homophenylalanine, citrulline, kynurenine, 4-aminophenylalanine, 3-(2-naphthyl)-alanine, 3-(1-naphthyl)alanine, methionine sulfone, t-butyl-alanine 5 t-butylglycine, 4-hydroxyphenylglycine, aminoalanine. phenylglycine, vinylalanine, propargyl-glycine, 1,2,4-triazolo-3-alanine, 4,4,4-trifluoro-threonine, thyronine, 6-hydroxytryptophan, 5-hydro-xytryptophan, 3-hydroxykynurenine, 3-aminotyrosine, trifuoromethyl-10 alanine, 2-thienylalanine, (2-(4-pyridyl)ethyl)cysteine, 3,4-dimethoxy-phenylalanine, 3-(2-thiazolyl)alanine, ibotenic acid, 1-amino-1-cyclopentanecarboxylic acid, 1-amino-1-cyclohexanecarboxylic acid, quisqualic acid, 3-trifluoromethylphenylalanine, 15 4-trifluoro-methylphenylalanine, cyclohexylalanine, cyclo-hexylglycine, thiohistidine, 3-methoxytyrosine. elastatinal, norleucine, norvaline, alloisoleucine. homoarginine, thioproline, dehydroproline, hydroxyproline, isonipectotic acid, homoproline, cyclohexyl-20 glycine, a-amino-n-butyric acid, cyclohexylalanine, aminophenylbutyric acid, phenylalanines substituted at the ortho, meta, or para position of the phenyl moiety with one or two of the following: a (C_1-C_4) alkyl, a (C_1-C_4) alkoxy, halogen or nitro groups or substituted 25 with a methylenedioxy group; β-2- and 3-thienylalanine, β -2- and 3-furanvlalanine, β -2-, 3- and 4-pyridylalanine, β -(benzothienyl-2- and 3-yl)alanine. β -(1- and 2-naphthyl)alanine, 0-alkylated derivatives of serine, threonine or tyrosine, S-alkylated cysteine, S-alkylated homocysteine, O-sulfate, O-phosphate and O-30 carboxylate esters of tyrosine, 3-sulfo-tyrosine, 3carboxy-tyrosine, 3-phospho-tyrosine, 4-methane sulfonic acid ester of tyrosine, 4-methane phosphonic acid ester of tyrosine, 3,5-diiodotyrosine, 3-nitro-

tyrosine, ε-alkyl lysine, and delta-alkyl ornithine.

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Any of these α -amino acids may be substituted with a methyl group at the alpha position, a halogen at any aromatic residue on the α -amino side chain, or an appropriate protective group at the O, N, or S atoms of the side chain residues. Appropriate protective groups are disclosed in "Protective Groups In Organic Synthesis," T.W. Greene and P.G.M. Wuts, J. Wiley & Sons, NY, NY, 1991.

The term "substitute" refers to the replacement of a hydrogen atom in a compound with a substituent group. In the present invention, those hydrogen atoms which form a part of a hydrogen bonding moiety which is capable of forming a hydrogen bond with the carbonyl oxygen of Arg-341 of ICE or the carbonyl oxygen of Ser-339 of ICE are excluded from substitution. These excluded hydrogen atoms include those which comprise an -NH- group which is alpha to a -CO- group and are depicted as -NH- rather than an X group or some other designation in the following diagrams: (a) through (t), (v) through (z).

The term "straight chain" refers to a contiguous unbranching string of covalently bound atoms. The straight chain may be substituted, but these substituents are not a part of the straight chain.

The term " K_1 " refers to a numerical measure of the effectiveness of a compound in inhibiting the activity of a target enzyme such as ICE. Lower values of K_1 reflect higher effectiveness. The K_1 value is a derived by fitting experimentally determined rate data to standard enzyme kinetic equations (see I. H. Segel, Enzyme Kinetics, Wiley-Interscience, 1975).

The term "patient" as used in this application refers to any mammal, especially numans.

The term "pharmaceutically effective amount" refers to an amount effective in treating or

- 21 -

ameliorating an IL-1-, apoptosis-, IGIF- or IFN-y-mediated disease in a patient. The term "prophylactically effective amount" refers to an amount effective in preventing or substantially lessening IL-1-, apoptosis-, IGIF or IFN-y mediated diseases in a

IL-1-, apoptosis-, IGIF or IFN-γ mediated diseases in a patient.

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The term "pharmaceutically acceptable carrier or adjuvant" refers to a non-toxic carrier or adjuvant that may be administered to a patient, together with a compound of this invention, and which does not destroy the pharmacological activity thereof.

The term "pharmaceutically acceptable derivative" means any pharmaceutically acceptable salt, ester, or salt of such ester, of a compound of this invention or any other compound which, upon administration to a recipient, is capable of providing (directly or indirectly) a compound of this invention or an anti-TCR active metabolite or residue thereof.

Pharmaceutically acceptable salts of the 20 compounds of this invention include, for example, those derived from pharmaceutically acceptable inorganic and organic acids and bases. Examples of suitable acids include hydrochloric, hydrobromic, sulfuric, nitric. perchloric, fumaric, maleic, phosphoric, glycolic, lactic, salicylic, succinic, toluene-p-sulfonic, 25 tartaric, acetic, citric, methanesulfonic, formic. benzoic, malonic, naphthalene-2-sulfonic and benzenesulfonic acids. Other acids, such as oxalic. while not in themselves pharmaceutically acceptable, may be employed in the preparation of salts useful as 30 intermediates in obtaining the compounds of the invention and their pharmaceutically acceptable acid addition salts. Salts derived from appropriate bases include alkali metal (e.g., sodium), alkaline earth

metal (e.g., magnesium), ammonium and N-(C1-4 alkyl),

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salts.

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This invention also envisions the "quaternization" of any basic nitrogen-containing groups of the compounds disclosed herein. The basic nitrogen can be quaternized with any agents known to those of ordinary skill in the art including, for example, lower alkyl halides, such as methyl, ethyl, propyl and butyl chloride, bromides and iodides; dialkyl sulfates including dimethyl, diethyl, dibutyl and diamyl sulfates; long chain halides such as decyl, lauryl, myristyl and stearyl chlorides, bromides and iodides; and aralkyl halides including benzyl and phenethyl bromides. Water or oil-soluble or dispersible products may be obtained by such quaternization.

The ICE inhibitors of this invention may contain one or more "asymmetric" carbon atoms and thus may occur as racemates and racemic mixtures, single enantiomers, diastereomeric mixtures and individual 20 diastereomers. All such isomeric forms of these compounds are expressly included in the present invention. Each stereogenic carbon may be of the R or S configuration. Although specific compounds and scaffolds exemplified in this application may be 25 depicted in a particular stereochemical configuration, compounds and scaffolds having either the opposite stereochemistry at any given chiral center or mixtures thereof are also envisioned. The ICE inhibitors of this invention may comprise ring structures which may optionally be 30 substituted at carbon, nitrogen or other atoms by various substituents. Such ring structures may be singly or multiply substituted. Preferably, the ring structures contain between 0 and 3 substituents. When

multiply substituted, each substituent may be picked

independently of any other substituent as long as the

- 23 -

combination of substituents results in the formation of a stable compound.

Combinations of substituents and variables envisioned by this invention are only those that result in the formation of stable compounds. The term "stable", as used herein, refers to compounds which possess stability sufficient to allow manufacture and administration to a mammal by methods known in the art. Typically, such compounds are stable at a temperature of 40°C or less, in the absence of moisture or other chemically reactive conditions, for at least a week.

Substituents may be represented in various forms. These various forms are known to the skilled practitioner and may be used interchangeably. For example, a methyl substituent on a phenyl ring may be represented in any of the following forms:

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Various forms of substituents such as methyl are used herein interchangeably.

DETAILED DESCRIPTION OF THE INVENTION

In order that the invention herein described may be more fully understood, the following detailed description is set forth.

The ICE inhibitors of one embodiment (A) of this invention are those of formula $\alpha\colon$

$$\begin{array}{c} (\text{CJ}_2)_m - \text{T} \\ \text{Q} \\ \text{R}_1 - \text{NH} - \text{X}_1 \\ (\text{CH}_2)_m - \text{R}_3 \end{array}$$

wherein:

- 24 -

$$X_1$$
 is -CH;

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q is 0 or 1;

each J is independently selected from the group consisting of -H, -OH, and -F, provided that when a first and second J are bound to a C and said first J is -OH, said second J is -H;

m is 0, 1, or 2;

T is -OH, -CO-CO₂H, -CO₂H, or any bioisosteric replacement for -CO₂H;

R₁ is selected from the group consisting of the following formulae, in which any ring may optionally be singly or multiply substituted at any carbon by Q₁, at any nitrogen by R₅, or at any atom by =0, -OH, -CO₂H, or halogen; any saturated ring may optionally be unsaturated at one or two bonds; and wherein R₁ (e) and

 R_1 (y) are optionally benzofused;

- 25 -

$$(d) \qquad \begin{matrix} X \\ \\ X \\ \\ N \end{matrix} \qquad \begin{matrix} R_6 \\ \\ -C \\ -C \\ \\ H \end{matrix} \qquad , \label{eq:controller}$$

- 28 -

 R_{20} is selected from the group consisting of:

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wherein each ring C is independently chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

$$\begin{array}{c} R_3 \text{ is:} \\ -\text{CN,} \\ -\text{CH=CH-R}_9, \\ -\text{CH=N-O-R}_9, \\ 15 & -(\text{CH}_2)_{1-3}\text{-T}_1\text{-R}_9, \\ -\text{CJ}_2\text{-R}_9, \\ -\text{CO-R}_{13}, \text{ or} \\ -\text{CO-CO-N} \\ R_{10}; \end{array}$$

each $\ensuremath{\mathtt{R}}_4$ is independently selected from the group consisting of:

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-(CH_2)_{1,2,3}-T_1-R_9;
```

each \mathbf{T}_1 is independently selected from the group consisting of:

CH=CH-, 5 -0-, -s-, -SO-, -SO2-, -NR10-, 10 -NR₁₀-CO-, -CO-, -0-CO-, -CO-O-, -CO-NR₁₀-, 15 -O-CO-NR₁₀-, -NR₁₀-CO-O-, -NR10-CO-NR10-, -SO2-NR10-, -NR10-SO2-, and -NR₁₀-SO₂-NR₁₀-; 20

each $R_{\mbox{\scriptsize 5}}$ is independently selected from the group consisting of:

-H,
-Ar₁,
-CO-Ar₁,
-SO₂-Ar₁,
-CO-NH₂,
-SO₂-NH₂,
-R₉,
30
-CO-R₉,

-CO-O-R₉,

- 32 -

$$\begin{array}{ccc} & & & /\text{Ar}_1 \\ & -\text{SO}_2\text{-N} \\ & \text{N}_{10}, \end{array}$$

-CO-N
$$R_{10}$$
, and

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 $R_{\rm 6}$ and $R_{\rm 7}$ taken together form a saturated 4-8 member carbocyclic ring or heterocyclic ring containing

-O-, -S-, or -NH-; or R_7 is -H and R_6 is -H

-Ar1,

-Rq,

 $-(CH_2)_{1,2,3}-T_1-R_9$, or

an α-amino acid side chain residue;

each R_9 is a C_{1-6} straight or branched alkyl group optionally singly or multiply substituted with -OH, -F, or =O and optionally substituted with one or two Ar_1 groups;

each R_{10} is independently selected from the group consisting of -H or a C_{1-6} straight or branched alkyl group;

each $R_{\rm 13}$ is independently selected from the group consisting of -Ar2, -R4 and -N-OH

each Ar; is a cyclic group independently selected

- 33 -

from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, a cycloalkyl group which contains between 3 and 15 carbon atoms and between 1 and 3 rings, said cycloalkyl group being optionally benzofused, and a 5 heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocycle group containing at least one heteroatom group selected from -O-, -S-, -SO-, -SO₂-, =N-, and -NH-, said heterocycle group optionally containing one or more double bonds. 10 said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted with -NH2, -CO2H, -Cl, -F, -Br, -I, -NO2, -CN, 15

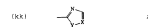
=0, -OH, -perfluoro
$$C_{1-3}$$
 alkyl, $\begin{pmatrix} O \\ \\ CH_2 \end{pmatrix}$, or $-Q_1$;

20 each Ar₂ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by -Q₁ and -Q₂:

(hh)
$$\stackrel{Y}{\longrightarrow}$$
 ; (ii) $\stackrel{X}{\longrightarrow}$; and

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each \mathbf{Q}_1 is independently selected from the group consisting of:

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each $\rm O_2$ is independently selected from the group consisting of -OH, -NH $_2$, -CO $_2$ H, -Cl, -F, -Br, -I, -NO $_2$, -CN, -CF $_3$, and O

provided that when $-Ar_1$ is substituted with a Q_1 group which comprises one or more additional $-Ar_1$ groups, said additional $-Ar_1$ groups are not substituted with Q_1 ;

each X is independently selected from the group consisting of =N-, and =CH-;

each X_2 is independently selected from the group consisting of -O-, -CH2-, -NH-, -S-, -SC-, and -SO2-;

each x_3 is independently selected from the group consisting of -CH2-, -S-, -SO-, and -SO2-;

each X_4 is independently selected from the group consisting of -CH $\!_2-$ and -NH- $\!_3$

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each X_5 is independently selected from the group
       consisting of -CH- and -N-;
             X6 is -CH- or -N-;
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             each Y is independently selected from the group
       consisting of -O-, -S-, and -NH;
             each Z is independently CO or SO2;
             each a is independently 0 or 1;
            each c is independently 1 or 2;
10
            each d is independently 0, 1, or 2; and
            each e is independently 0, 1, 2, or 3;
       provided that when
                  R_1 is (f),
15
                  R_6 is an \alpha-amino acid side chain residue, and
                  R7 is -H,
            then (aa1) and (aa2) must be substituted with Q_1;
            also provided that when
20
                 R; is (o),
                 q is 0.
                 J is -H.
                 m is 1,
                 R_6 is an \alpha-amino acid side chain residue.
25
                 Ry is -H.
                 X2 is -CH2-,
                 Xs is -CH- ,
                 X_6 is -N- , and
30
                           -CO-N _{\rm R_{10}}^{\rm R_{10}} , or -CO-R_{\rm 13}, when
                 R_3 is
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R₁₃ is: -CH₂-O-CO-Ar₁, -CH₂-S-CO-Ar₁, -CH₂-O-Ar₁, -CH₂-S-Ar₁, or -R₄ when -R₄ is -H₁

then the ring of the $R_1(o)$ group must be substituted with Q_1 or benzofused; and

provided that when

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 R_1 is (w),

g is 0,

J is -H,

m is 1, T is -CO₂H,

1 18 -CO

 X_2 is O,

 R_5 is benzyloxycarbonyl, and

ring C is benzo,

then R_3 cannot be -CO- R_{13} when:

 R_{13} is -CH₂-O-Ar₁ and

Ar₁ is 1-phenyl-3-trifluoromethylpyrazole-5-yl wherein the phenyl is optionally substituted with a chlorine atom:

or when

 ${\tt R}_{13}$ is -CH $_2$ -O-CO-Ar $_1$, wherein

 Ar_1 is 2,6-dichlorophenyl.

 $\label{eq:preferred compounds of embodiment A employ}$ formula $\alpha,$ wherein R_1 is (w):

(w) R₀ N C C C H

wherein the other substituents are as described

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above.

Other preferred compounds of embodiment A employ formula $\alpha,$ wherein R_1 is $(y)\colon$

$$(y) \begin{array}{c} x_{T^{*}}(CH_{2})_{c} \\ (CH_{1})_{n} \\ x_{5}^{*}(CH_{2})_{c} \\ X_{3} \\ (CH_{2})_{c} \\ \end{array}$$

wherein the other substituents are as described above.

 $\label{eq:more preferred compounds of embodiment A} \text{ employ formula } \alpha, \text{ wherein:}$

$$X_1$$
 is -CH;

10 q is 0;

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J is -H;

m is 0 or 1 and T is -CO-CO $_2\mathrm{H},$ or any bioisosteric replacement for -CO $_2\mathrm{H},$ or

m is 1 and T is -CO2H;

 R_1 is selected from the group consisting of the following formulae, in which any ring may optionally be singly or multiply substituted at any carbon by Q_1 , at any nitrogen by R_5 , or at any atom by =0, -0H, -CO₂H, or halogen, and wherein (e) is optionally benzofused:

$$\begin{array}{c} \text{(c)} \\ \text{Rs} \\ \text{N} \\ \text{H} \\ \text{O} \end{array}$$

$$(x) \qquad \begin{array}{c} (x) \\ X_{2} \\ X_{3} \\ X_{4} \\ X_{5} \\ X_{5} \\ X_{7} \\ X_{7} \\ X_{7} \\ X_{8} \\ X_{9} \\ X_{9} \\ X_{9} \\ X_{1} \\ X_{1} \\ X_{2} \\ X_{3} \\ X_{4} \\ X_{1} \\ X_{2} \\ X_{3} \\ X_{4} \\ X_{1} \\ X_{2} \\ X_{3} \\ X_{4} \\ X_{5} \\ X_{5} \\ X_{5} \\ X_{7} \\ X_{1} \\ X_{2} \\ X_{3} \\ X_{4} \\ X_{5} \\ X_{$$

ring C is benzo optionally substituted with $\label{eq:condition} \mbox{-C1}_{1-3} \mbox{ alkyl, -O-C1}_{1-3} \mbox{ alkyl, -Cl, -F or -CF3},$

when R_1 is (a) or (b), R_5 is preferably -H, and

when R_1 is (c), (e), (f), (o), (r), (w), (x) or $\label{eq:state} (y) \;,\; R_5 \; \text{is preferably};$

and c is 1;

20 -CO-R₉, -CO-O-R₉.

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$$\ensuremath{\mathtt{R}}_7$$
 is -H and $\ensuremath{\mathtt{R}}_6$ is: -H,
$$-\ensuremath{\mathtt{R}}_9, \text{ or } \\ -\ensuremath{\mathtt{Ar}}_1;$$

 $R_{\rm g}$ is a C_{1-6} straight or branched alkyl group optionally substituted with =0 and optionally substituted with -Ar₁;

 $$\rm R_{10}$ is -H or a -C $_{\rm 1-3}$ straight or branched alkyl $$\rm group;$

Ar_1 is phenyl, naphthyl, pyridyl, benzothiazolyl, thienyl, benzothienyl, benzoxazolyl, 2-indanyl, or indolyl optionally substituted with -O- C_{1-3} alkyl, -N- C_{1-3} alkyl, -N- C_{1-3} alkyl, -N- C_{1-3} alkyl)₂, -Cl, -F, -CF₃,

Q₁ is R₉ or $-(CH_2)_{0,1,2}-T_1-(CH_2)_{0,1,2}-Ar_1$, wherein T_1 is -O- or -S-;

each X is independently selected from the group consisting of =N-, and =CH-;

each x_2 is independently selected from the group consisting of -O-, -CH₂-, -NH-, -S-, -SO-, and -SO₂-;

each $\rm X_5$ is independently selected from the group consisting of -CH- and -N-; |

$$X_6$$
 is -CH- or -N-,

- 41 -

provided that when:

 R_1 is (o),

 X_2 is -CH₂-,

 X_5 is -CH- , and

 X_6 is -N- ,

then the ring of the $R_1(o)$ group must be substituted with ${\bf Q}_1$ or benzofused; and

Z is C=O.

Most preferably, compounds of this more preferred embodiment are those wherein the $\rm R_1$ group is:

(e1)

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Re N

, or

(e2)

and c is 2; or

(e4)

, or

- 42 -

which is optionally benzofused, and c is 1 or 2;

provided that when R_1 is (e4),

5 g is 0,

J is -H,

m is 1,

T is $-CO_2H$,

R₅ is benzyloxycarbonyl, and

10 c is 1,

then R_3 cannot be -CO- R_{13} when

 R_{13} is $-CH_2-O-Ar_1$ and

Ar₁ is 1-phenyl-3-trifluoromethyl-pyrazole-

5-yl, wherein the phenyl is optionally substituted with 15 a chlorine atom; or when

 R_{13} is -CH $_2$ -O-CO-Ar $_1$, wherein

Ar₁ is 2,6-dichlorophenyl,

and when the 2-position of the scaffold ring is

20 substituted with para-fluoro-phenyl; and

also provided that when

 R_1 is (e7),

q is 0,

J is -H,

m is 1.

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T is CO2H or -CO-NH-OH,

 R_{\S} is a protective group for the N atom of an amino acid side chain residue, and

each c is 1.

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then
$$R_3$$
 cannot be $-CO-R_{13}$ when R_{13} is: $-CH_2-O-CO-Ar_1$, $-CH_2-S-CO-Ar_1$, $-CH_2-O-Ar_1$, or $-CH_2-S-Ar_1$.

The most preferred compounds of this embodiment are those wherein:

R₁ is:

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and c is 2;

m is 1; T is -CO₂H; and
$$R_3$$
 is -CO- R_{13} .

Other most preferred compounds of this embodiment are those wherein:

R₁ is:

optionally substituted with R_5 or Q_1 at X_2 when X_2 10 $\,$ is -NH-; and

ring C is benzo substituted with ${\rm -C_{1-3}}$ alkyl, ${\rm -O-C_{1-3}}$ alkyl, -Cl, -F or -CF3.

The ICE inhibitors of another embodiment (B) of this invention are those of formula (I):

-SO₂-, or -NH-;

wherein:

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 $$R_{1}$$ is selected from the group consisting of the 20 $\,$ following formulae:

$$\begin{array}{c} \text{(el0)} \\ \text{R}_{2}\text{-} \\ \text{R}_{5}\text{-} \\ \text{H} \end{array}$$

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(e11)
$$R_{3} - N \qquad ;$$

$$R_{2} - N \qquad ;$$

$$R_{3} - N \qquad ;$$

$$R_{5} - N \qquad ;$$

$$R_{5}$$

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

- 46 -

 R_2 is:

m is 1 or 2;

 X_7 is $-N(R_8)$ - or -0-;

20
$$X_5$$
 is -CH- or -N-; Y_2 is H_2 or O;

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 $R_{\rm 6}$ is selected from the group consisting of -H and -CH $_{\rm 3};$

```
 R_8 \text{ is selected from the group consisting of:} \\  -C(0)-R_{10}, \\ -C(0)-R_9, \\ -C(0)-N(H)-R_{10}, \\ -S(0)_2-R_9, \\ -S(0)_2-NH-R_{10}, \\ 10 & -C(0)-CH_2-OR_{10}, \\ -C(0)-CH_2-OR_{10}, \\ -C(0)-CH_2N(R_{10})(R_{10}), \\ -C(0)-CH_2C(0)-O-R_9, \\ -C(0)-CH_2C(0)-O-R_9, \\ 15 & -H, \text{ and } \\ -C(0)-C(0)-C(0)-OR_{10}, \\ \end{aligned}
```

each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

20

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each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 R_{13} is selected from the group consisting of H, Ar3, and a C_{1-6} straight or branched alkyl group optionally substituted with Ar3, -CONH2, -OR5, -OH, -OR9, or -CO2H;

each R_{51} is independently selected from the group consisting of R_{9} , -C(O)- R_{9} , -C(O)-N(H)- R_{9} , or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

each R_{21} is independently selected from the group consisting of -H or a - C_{1-6} straight or branched alkyl group;

each Ar₃ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

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each Q_1 is independently selected from the group consisting of -NH₂, -CQ₂H₁, -Cl, -F, -Br, -I, -NO₂, -CN, =0, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, OR₉, -NHR₀, R_0 , -C(0)- R_{10} , and

30

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

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Preferably, R_5 is selected from the group consisting of: $-C(0) - R_{10}, \\ -C(0) O - R_9, \text{ and } \\ -C(0) - NH - R_{10}.$

 $\label{eq:alternatively, R5} \mbox{ Alternatively, R$_5$ is selected from the group consisting of:}$

 $-S(0)_2-R_9$, $-S(0)_2-NH-R_{10}$, $-C(0)-C(0)-R_{10}$,

 $-R_9$, and $-C(0) - C(0) - OR_{10}$.

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More preferably:

m is 1;

R₁₃ is H or a -C₁₋₄ straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, or -CO₂H, wherein the R₉ is a -C₁₋₄ branched or straight alkyl group, wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O₁;

20 R₂₁ is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar3, wherein Ar3 is phenyl, optionally substituted by $-Q_1\,;$

25 Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl benzofuranyl, and indolyl;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -NHR₉, and

wherein each R₉ and R₁₀ are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃ wherein Ar₃ is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

The ICE inhibitors of another embodiment (C) of this invention are those of formula (\underline{II}):

wherein:

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m is 1 or 2;

 $\ensuremath{R_{\mathrm{1}}}$ is selected from the group consisting of the following formulae:

(e11)
$$R_{S}-N$$

oring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

; and

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R_3 is selected from the group consisting of:
                   -CN,
                   -C(O)-H,
                   -C(0)-CH2-T1-R11,
                   -C(0)-CH2-F,
 5
                   -C=N-O-Rg, and
                   -CO-Ar2;
             Rs is selected from the group consisting of:
                   -C(O)-R<sub>10</sub>,
                   -C(0)0-Rg,
10
                   -C(0)-N \begin{pmatrix} R_{10} \\ R_{10} \end{pmatrix}
15
                   -S(0)2-Rg,
                   -C(0)-CH2-O-R9,
                   -C(0)C(0)-R10.
20
                   -R9.
                   -H, and
                   -C(0)C(0)-OR10.
             x_5 is -CH- or -N-;
25
             Y2 is H2 or O;
             X_7 is -N(R_8) - or -O-;
             each T1 is independently selected from the group
       consisting of -O-, -S-, -S(O)-, and -S(O)2-;
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```

Rg is selected from the group consisting of:

- CH3;

Re is selected from the group consisting of -H and

- 53 -

 $\begin{array}{c} -C(O) - R_{10}, \\ -C(O) - R_{9}, \\ -C(O) - NH - R_{10}, \\ -S(O) _{2} - R_{9}, \\ \end{array}$ $5 \qquad -S(O) _{2} - NH - R_{10}, \\ -C(O) - CH_{2} - OR_{10}, \\ -C(O) - CH_{2} - N(R_{10}) (R_{10}), \\ -C(O) - CH_{2} - CH_{20}(O) - OR_{9}, \\ -C(O) - CH_{2} - C(O) - CH_{20}(O) - R_{10}, \\ -C(O) - CH_{20}(O) - CH_{20}(O) - R_{10}, \\ -C(O) - CH_{20}(O) - CH_{20}(O) - R_{10}, \\ -C(O) - C(O) -$

each R_9 is independently selected from the group consisting of -Ar₃ and a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each \mathbf{R}_{11} is independently selected from the group consisting of:

-Ar₄, -(CH₂)₁₋₃-Ar₄, -H, and -C(0)-Ar₄;

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 R_{13} is selected from the group consisting of H, Ar₃, and a C_{1-6} straight or branched alkyl group optionally substituted with Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

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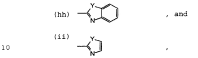
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-OR-3 is optionally -N(H)-OH;

each ${\rm R}_{21}$ is independently selected from the group consisting of -H or a -C $_{1-6}$ straight or branched alkyl group;

Ar₂ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by -Q₁:



wherein each Y is independently selected from the group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, -N(R₃)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each ${\rm Ar_4}$ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and

- 55 -

15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -0-, -S-, -SO-, SO₂, -N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -01;

each Q_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =O, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, OR₉, -NHR₉, R_9 , -C(O)-R₁₀, and

provided that when -Ar $_3$ is substituted with a Q $_1$ group which comprises one or more additional -Ar $_3$ with another -Ar $_3$.

Preferred compounds of this embodiment include, but are not limited to:

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- 57 -

- 58 -

Preferred compounds of embodiment C employ formula (II), wherein \mathbf{R}_1 is (e11) and the other substituents are as defined above.

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Other preferred compounds of embodiment C employ formula (II), wherein \mathbf{R}_1 is (e12) and the other substituents are as defined above.

Other preferred compounds of embodiment C employ formula (II) wherein R_1 is (y1) and the other substituents are as defined above.

Other preferred compounds of embodiment C employ formula (II) wherein \mathbf{R}_1 is (y2) and the other substituents are as defined above.

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Other preferred compounds of embodiment C of employ formula (II) wherein R_1 is (z) and the other substituents are as defined above.

Other preferred compound of embodiment C $\,$ employ formula (II) wherein R_1 is (w2) and the other substituents are as defined above.

More preferably, R1 is (w2) and

m is 1:

ring C is benzo, pyrido, or thieno;

 $\ensuremath{R_{5}}$ is selected from the group consisting of:

-C(O)- R_{10} , wherein R_{10} is -Ar₃;

-C(0)0-R₉, wherein R₉ is -CH₂-Ar₃;

-C(O)C(O)-R $_{10}$, wherein R $_{10}$ is -CH $_{2}$ Ar $_{3}$;

 $^{-R}{\rm g},$ wherein ${\rm R}_{9}$ is a ${\rm C}_{1-2}$ alkyl group substituted with $^{-A}{\rm r}_{3};$ and

-C(0)C(0)-OR₁₀, wherein R₁₀ is -CH₂Ar₃;

20 T₁ is 0 or S;

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R6 is H;

 R_{8} is selected from the group consisting -C(0)-R₁₀, -C(0)-CH₂-OR₁₀, and -C(0)CH₂-N(R₁₀)(R₁₀), wherein R₁₀ is H, CH₃, or -CH₂CH₃;

25 R_{11} is selected from the group consisting of -Ar₄, -(CH₂)₁₋₃-Ar₄, and -C(O)-Ar₄;

- 60 -

 R_{13} is H or a $-C_{1-4}$ straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, or $-CO_2H$, wherein the R_9 is a $-C_{1-4}$ branched or straight alkyl group, wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

Ar2 is (hh);

Y is 0;

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Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl, thienothienyl, thiadiazolyl, benzotriazolyl, benzo[b]thiophenyl, benzofuranyl, and indolyl;

Ar₄ is phenyl, tetrazolyl, naphthyl, pyridinyl, oxazolyl, pyrimidinyl, or indolyl;

each Q_1 is independently selected from the group consisting of "NH2, -Cl, -F, -Br, -OH, -R9, -NH-R5 wherein R_5 is -C(0)-R10 or -S(0)₂-R9, -OR5 wherein R_5 is -C(0)-R10, -OR4, -NHR4, and

CH₂

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_4$.

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Preferred compounds of this embodiment include, but are not limited to:

- 62 -

- 64 -

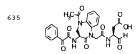
- 65 -

- 66 -

627

- 67 -

- 68 -



Other preferred compounds of embodiment C employ formula (II) wherein R_1 is (e10), X_5 is CH, and the other substituents are as defined above.

More preferred compounds of embodiment C employ formula (II) wherein R_1 is (e10), X_5 is CH, R_3 is CO-Ar₂, and the other substituents are as defined above.

Other more preferred compounds of embodiment C employ formula (II) wherein R_1 is (e10), X_5 is CH, R_3 is -C(O)-CH₂-T₁-R₁₁, R_{11} is -(CH₂)₁₋₃-Ar₄, and the other substituents are as defined above.

Other more preferred compounds of embodiment C employ formula (II) wherein \Re_1 is (e10) and \aleph_5 is CH and

$$R_3$$
 is $-C(O) - CH_2 - T_1 - R_{11}$;
 T_1 is O ; and
 R_{11} is $-C(O) - Ar_4$,

20 and the other substituents are as defined above.

More preferably, in these more preferred compounds, R_5 is selected from the group consisting of:

- -C(O)-R10,
- -C(0)0-Rq, and
- 25 -C(0)-NH-R₁₀.

5

1.0

- 69 -

Alternatively, in these more preferred compounds, R_5 is selected from the group consisting of:

 $-S(0)_2-R_9$, $-S(0)_2-NH-R_{10}$, $-C(0)-C(0)-R_{10}$, $-R_9$, and $-C(0)-C(0)-OR_{10}$.

Most preferably, in these more preferred compounds,

m is 1;

T₁ is O or S;

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 R_{13} is H or a $-C_{1-4}$ straight or branched alkyl group optionally substituted with $-Ar_3$, -OR, $-OR_9$, or $-CO_2H$, wherein the R_9 is a $-C_{1-4}$ branched or straight alkyl group, wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

 R_{21} is -H or -CH3;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl, optionally substituted by $-Q_1$;

Aro is (hh);

Y is O, and

Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl benzofuranyl, and indolyl;

- 70 -

Ar₄ is phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₈, and

CH₂

wherein each R_9 and R_{10} are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

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3.0

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1.0

provided that when -Ar $_3$ is substituted with a Q $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Other more preferred compounds of embodiment C employ formula (II) wherein R_1 is (e10), X_5 is CH, R_3 is -C(0)-H, and the other substituents are as defined above.

More preferably, in these more preferred compounds, R_5 is selected from the group consisting of:

Alternatively, in these more preferred compounds, R_5 is selected from the group consisting of:

 $⁻s(0)_2-NH-R_{10}$,

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 $-C(0)-C(0)-R_{10}$,

-Rq, and

-C(O)-C(O)-OR10.

Most preferably, in these more preferred compounds,

5 m is 1;

15

 T_1 is 0 or S;

 R_{13} is H or a $-C_{1-4}$ straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, or 10 $-CO_2H$, wherein the R_9 is a $-C_{1-4}$ branched or straight alkyl group, wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

 R_{21} is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl, optionally substituted by $-Q_1$;

 Ar_2 is (hh);

Y is O, and

20 Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl benzofuranyl, and indolyl;

25 Ar₄ is phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl;

- 72 -

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -NHR₉, and

O /\CH₂,

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wherein each R₉ and R₁₀ are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃ wherein Ar₃ is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$,

Other more preferred compounds of embodiment C employ formula (II) wherein R_1 is (e10) and X_5 is CH, R_3 is -CO-CH₂-T₁-R₁₁, and R_{11} is -Ar₄, and the other substituents are as defined above.

More preferably, in these more preferred compounds, $\rm R_{\rm 5}$ is selected from the group consisting of:

- -C(O)-R10,
- -C(0)0-Rq, and
- -C(0)-NH-R10.

Alternatively, in these more preferred compounds, \Re_5 is selected from the group consisting of:

-S(0)2-Rg,

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 $-S(O)_2-NH-R_{10}$, $-C(O)_-C(O)_-R_{10}$, $-R_9$, and $-C(O)_-C(O)_-OR_{10}$.

5 Most preferably, in these more preferred compounds,

m is 1;

 T_1 is 0 or S;

R₁₃ is H or a -C₁₋₄ straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, or -CO₂H, wherein the R₉ is a -C₁₋₄ branched or straight alkyl group, wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q₁;

 R_{21} is -H or -CH₃;

15 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl, optionally substituted by $-Q_1$;

Aro is (hh);

Y is O, and

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Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl benzofuranyl, and indolyl:

Ar₄ is phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl;

- 74 -

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₉, and

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O CH₂,

wherein each R₉ and R₁₀ are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃ wherein Ar₃ is phenyl;

provided that when -Ar $_3$ is substituted with a Q $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Other preferred compounds of embodiment C employ formula (II) wherein R_1 is (e10), X_5 is N, and the other substituents are as defined above.

More preferred compounds of embodiment C, employ formula (II) wherein R_1 is (e10), X_5 is N, R_3 is CO-Ar₂, and the other substituents are as defined above.

Other more preferred compounds of embodiment C, employ formula (II) wherein R_1 is (e10), X_5 is N, R_3 is -C(0)- CH_2 - T_1 - R_{11} , R_{11} is $-(CH_2)_{1-3}$ - Ar_4 , and the other substituents are as defined above.

Other more preferred compounds of embodiment C, employ formula (II) wherein R1 is (e10) and $\rm X_5$ is N and:

- 75 -

$$R_3$$
 is $-C(0) - CH_2 - T_1 - R_{11}$;
 T_1 is 0; and

 $\ensuremath{\mathtt{R}}_{11}$ is -C(O)-Ar4, and the other substituents are as defined above.

5 More preferably, in these more preferred compounds, R_5 is selected from the group consisting of:

$$-C(0)-R_{10}$$
,
 $-C(0)0-R_{9}$, and

-C(O)-NH-R₁₀.

Alternatively, in these more preferred compounds, R₅ is selected from the group consisting of:

$$-S(0)_2-R_9$$
,
 $-S(0)_2-NH-R_{10}$,
 $-C(0)-C(0)-R_{10}$,

-R₉, and -C(O)-C(O)-OR₁₀.

1.5

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Most preferably, in these more preferred compounds, $R_{\bar{\bf 5}}$ is selected from the group consisting of:

m is 1;

 T_1 is 0 or S;

 R_{13} is H or a $-C_{1-4}$ straight or branched alkyl group optionally substituted with $-Ar_{3}$, -OH, $-OR_{9}$, or $-CO_{2}H$, wherein the R_{9} is a $-C_{1-4}$ branched or straight alkyl group, wherein Ar_{3} is morpholinyl or phenyl, wherein the phenyl is optionally substituted with $Q_{1,7}$

- 76 -

R21 is -H or -CH3;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl, optionally substituted by -0:

5 Ar₂ is (hh);

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Y is O, and

Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo(b)thiophenyl, pyridyl benzofuranyl, and indolyl;

Ar₄ is phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₄, and

O /\CH₂,

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$ wherein Ar $_3$ is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, sa. additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

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Other more preferred compounds of embodiment C, employ formula (II) wherein R_1 is (ei0), X_5 is N, R_3 is -C(0)-H, and the other substituents are as defined above.

5 More preferably, in these more preferred compounds, R_5 is selected from the group consisting of:

-C(O)-R10.

-C(0)0-R9, and

 $-C(0)-NH-R_{10}$.

Alternatively, in these more preferred compounds, R₅ is selected from the group consisting of:

-S(0)2-Rg,

 $-s(0)_2-NH-R_{10}$,

-C(0)-C(0)-R10,

-Ra, and

15

-C(O)-C(O)-OR₁₀.

Most preferably, in these more preferred compounds,

m is 1:

20 T₁ is 0 or S;

 R_{13} is H or a $^{-}C_{1-4}$ straight or branched alkyl group optionally substituted with $^{-}Ar_{3,}$ $^{-}OH,$ $^{-}OR_{9},$ or $^{-}CO_{2}H,$ wherein the R_{9} is a $^{-}C_{1-4}$ branched or straight alkyl group, wherein Ar $_{3}$ is morpholinyl or phenyl,

25 wherein the phenyl is optionally substituted with Q;

$$R_{21}$$
 is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl, optionally substituted by $-Q_7$;

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Aro is (hh):

Y is O. and

Ar₃ is phenyl, naphthyl, thienyl, quinolinyl,
isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl,
benzotriazolyl, benzimidazolyl, thienothienyl,
imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl
benzofuranyl, and indolyl;

Ar4 is phenyl, tetrazolyl, pyridinyl, oxazolyl,
10 naphthyl, pyrimidinyl, or thienyl;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -NHR₉, and

O /\CH₂,

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wherein each R₉ and R₁₀ are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃ wherein Ar₃ is phenyl;

provided that when -Ar $_3$ is substituted with a O $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Other more preferred compounds of embodiment C, employ formula (II) wherein R_1 is (e10), X_5 is N, R_3 is -CO-CH₂-T₁-R₁₁, R_{11} is -Ar₄, and the other substituents are as defined above.

- 79 -

More preferably, in these more preferred compounds, R_{5} is selected from the group consisting of:

-C(O)-R10,

-C(0)0-Ra, and

5 -C(0)-NH-R₁₀.

Alternatively, in these more preferred compounds, $\rm R_{\bar 5}$ is selected from the group consisting of:

-S(0)2-Rq,

-S(0)2-NH-R10,

-C(O)-C(O)-R₁₀,

-R₉, and

-C(O)-C(O)-OR₁₀.

Most preferably, in these more preferred compounds

m is 1;

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 T_1 is 0 or S;

 R_{13} is H or a $-C_{1-4}$ straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, or $-CO_2H$, wherein the R_9 is a $-C_{1-4}$ branched or straight alkyl group, wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl, optionally substituted by $-Q_1$;

Aro is (hh);

Y is O. and

- 80 -

Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl benzofuranyl, and indolyl;

Ar₄ is phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -NHR₉, and

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2.5

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

 $\label{eq:preferred} \mbox{ Preferred compounds of embodiment B include,} \\ \mbox{but are not limited to:}$

 $\label{eq:preferred compounds of embodiment C include,} \\ \text{but are not limited to:}$

- 82 -

- 83 -

- 84 -

- 85 -

- 86 -

418 PN N ON N ON H

- 87 -

- 88 -

- 90 -

- 91 -

- 92 -

- 91 -

- 95 -

- 96 -

- 97 -

- 98 -

- 99 -

- 100 -

- 101 -

5

817d

- 102 -

- 103 -

- 104 -

- 106 -

- 107 -

- 108 -

- 109 -

- 110 -

- 111 -

- 112 -

5 1054 ON HOOM H

- 114 -

- 115 -

- 116 -

- 117 -

- 118 -

- 119 -

- 121 -

- 122 -

Б

Specific compounds of this invention also include, but are not limited to, those compounds whose structures comprise scaffolds 1-22:

11
$$R_{S}$$
 R_{S}
 R

5

- 125 -

wherein:

5

.CO₂H

и сно

, wherein H
$$_{\rm H}$$
 $_{\rm CO_2R_{13}}$

- 126 -

each R_{51} is $-CH_3$, $-CH_2CH_3$, $-CH_2CH_2CH_3$, $-CH(CH_3)$ (CH_3), $-CH_2CH_2CH_2$, $-CH_2$ -CH(CH_3) (CH_3), $-CH_2$ -CH(CH_3) CH_3 , $-CH_3$ -CH(CH_3) CH_3 , $-CH_3$ -CH(CH_3) acetal or a propylenedioxy acetal; or

5

1.0

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25

 R_5 in each of the above compounds is the same as any one of the R_5 moieties shown for any one of compounds 139, 214c, 214e, 404-413, 415-491, 493-501.

Specific compounds of this invention also include, but are not limited to, compounds comprising scaffolds 1-28, wherein R, R_{51} , and R_{5} are as defined above, and in which the -C(O) - of the R_{5} moiety of any one of compounds 214c, 214e, 404-413, 415-418, 422-426, 430-456, 458-466, 468, 470-471, 473-491, 493, 495, 497-501 is replaced with -CH₂-, -C(O)C(O)-, or -CH₂C(O)C(O)-.

The ICE inhibitors of another embodiment (D) of this invention are those of formula (\underline{I}):

wherein:

 $\ensuremath{\mathtt{R}}_1$ is selected from the group consisting of the following formulae:

$$\begin{array}{c} R_8 \\ \\ R_5 - N \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\$$

- 128 -

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

R2 is:

m is 1 or 2;

 $\mbox{ each R_{\S} is independently selected from the group } \\ 15 & \mbox{ consisting of:} \\$

$$-C(0)-N(R_{10})(R_{10})$$

-S(O)2-R9,

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```
-S(O)2-NH-R10,
                   -C(O)-CH2-O-Rq,
                   -C(0)C(0)-R10
                  -Ra,
 5
                  -H,
                  -C(0)C(0)-OR10, and
                  -C(O)C(O)-N(Ra)(R10);
             X_5 is -CH- or -N-;
10
            Y_2 is H_2 or O;
            X_7 is -N(R_8) - or -O-;
             R6 is selected from the group consisting of -H and
15
       - CH3;
            R<sub>8</sub> is selected from the group consisting of:
                  -C(O)-R10,
                  -C(0)0-Rq,
                  -C(O)-N(H)-R10,
20
                  -S(O)2-Rq,
                  -S(0)2-NH-R10,
                  -C(O)-CH2-OR10,
                  -C(0)C(0)-R10;
25
                  -C(0) - CH_2N(R_{10})(R_{10}),
                  -C(O)-CH2C(O)-O-Rq,
                  -C(O)-CH2C(O)-R9,
                 -H, and
                  -C(0)-C(0)-OR10;
```

each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein

- 130 -

the -C1-6 alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 $\rm R_{13}$ is selected from the group consisting of H, Ar3, and a $\rm C_{1-6}$ straight or branched alkyl group optionally substituted with Ar3, -CONH2, -OR5, -OH, -OR9, or -CO2H;

each R_{51} is independently selected from the group consisting of R_9 , $-C(0)-R_9$, $-C(0)-N(H)-R_9$, or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

each R_{21} is independently selected from the group consisting of -H or a $-C_{1-6}$ straight or branched alkyl group;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁:

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1.0

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- 131 -

consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =O, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, OR₉, -N(R₉) (R_{10}), R_9 , -C(O)-R_{1C}, and

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provided that when -Ar $_3$ is substituted with a Q $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Preferably, R_5 is selected from the group

consisting of:

-C(O)-R₁₀,

 $-C(0)0-R_9$, and

-C(O)-NH- R_{10} .

Alternatively, $R_{\bar{\mathbf{5}}}$ is selected from the group consisting of:

-S(0)2-Rg,

-S(0)2-NH-R10,

-C(0)-C(0)-R10,

-Rq, and

25 -C(0)-C(0)-OR₁₀.

More preferably:

m is 1;

 R_{13} is H or a -C₁₋₄ straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, or -CO₂H, wherein the R_9 is a -C₁₋₄ branched or straight alkyl group, wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with $Q_1;$

- 132 -

R21 is -H or -CH3;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar3, wherein Ar3 is phenyl, optionally substituted by -Q1;

5

1.0

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each Ar3 cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q1;

each Q1 is independently selected from the group consisting of -NH2, -Cl, -F, -Br, -OH, -Rg, -NH-Rg wherein R_5 is $-C(0)-R_{10}$ or $-S(0)_2-R_9$, $-OR_5$ wherein R_5 is -C(O)-R10, -OR9, -N(R9)(R10), and

20

wherein each $\ensuremath{R_9}$ and $\ensuremath{R_{10}}$ are independently a $\ensuremath{\text{-C}_{1\text{--}6}}$ straight or branched alkyl group optionally substituted with Ara wherein Ara is phenyl:

provided that when -Ara is substituted with a Q. group which comprises one or more additional -Ara groups, said additional -Ara groups are not substituted with another -Ara.

The ICE inhibitors of another embodiment (E) of this invention are those of formula (II):

- 133 -

wherein:

5

m is 1 or 2;

 $\ensuremath{\mathtt{R}}_1$ is selected from the group consisting of the following formulae:

(e10)
$$\begin{array}{c} R_{2} \\ R_{5} - N \\ R_{5} - N \\ \end{array}$$
(e11)
$$\begin{array}{c} Y_{2} \\ N \\ R_{5} - N \\ \end{array}$$
(e12)
$$\begin{array}{c} Y_{2} \\ R_{5} - N \\ \end{array}$$

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$$(y1) \qquad \begin{array}{c} R_{8} & y_{2} \\ N & N \\$$

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

R₃ is selected from the group consisting of: -CN, -C(O)-H, -C(O)-CH₂- T_1 - R_{11} , -C(O)-CH₂-F, -C=N-O-R₉, and -CO-Ar₂;

each R_{5} is independently selected from the group consisting of:

20
$$-C(O) - R_{10}$$
,
 $-C(O) O - R_{9}$,
 $-C(O) - N(R_{10}) (R_{10})$
 $-S(O)_2 - R_{9}$,
 $-S(O)_2 - NH - R_{10}$,

5

- 135 -

each T_1 is independently selected from the group consisting of -O-, -S-, -S(O)-, and -S(O)2-;

15 $$R_{6}$$ is selected from the group consisting of -H and -CH3;

-C(0)-C(0)-OR10;

3.0

each R_{θ} is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein

- 136 -

the -C1-6 alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each $\ensuremath{R_{11}}$ is independently selected from the group consisting of:

-Ar₄, -(CH₂)₁₋₃-Ar₄, -H, and -C(O)-Ar₄;

5

15

 R_{15} is selected from the group consisting of -OH, -OAr₃, -N(H)-OH, and a -OC₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₄, or -CO₅H;

each $\rm R_{21}$ is independently selected from the group consisting of -H or a -C $_{1-6}$ straight or branched alkyl group;

Ar₂ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by -Q₁:

$$(hh)$$
 , and





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1.0

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3.0

wherein each Y is independently selected from the group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one hetercatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁:

each Ar_4 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

each Q_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, -O, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, OR₅, -N(R_9) (R_{10}), R_9 , -C(O)- R_{10} , and

- 138 -

1.0

1.5

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provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_4$.

Preferred compounds of embodiment E employ formula (II), wherein R_1 is (e11) and the other substituents are as defined above.

Other preferred compounds of embodiment E employ formula (II), wherein R_1 is (e12) and the other substituents are as defined above.

Other preferred compounds of embodiment E employ formula (II) wherein R_1 is (y1) and the other substituents are as defined above

Other preferred compounds of embodiment Ξ employ formula (II) wherein R_1 is (y2) and the other substituents are as defined above.

Other preferred compounds of embodiment E of employ formula (II) wherein R_1 is (z) and the other substituents are as defined above.

Other preferred compound of embodiment E employ formula (II) wherein R_1 is (w2) and the other substituents are as defined above.

More preferably, R1 is (w2) and

- 139 -

m is 1;

ring C is benzo, pyrido, or thieno;

 R_3 is selected from the group consisting of -C(O)-H, -C(O)-Ar2, and -C(O)CH2-T1-R11;

5 R₅ is selected from the group consisting of: -C(0)-R₁₀, wherein R₁₀ is -Ar₃; -C(0)O-R₉, wherein R₉ is -CH₂-Ar₃; -C(0)C(0)-R₁₀, wherein R₁₀ is -Ar₃; -R₄, wherein R₄ is a C₁₋₂ alkyl group

substituted with -Ar₃; and -C(0)C(0)-OR₁₀, wherein R_{10} is -CH₂Ar₃;

 T_1 is 0 or S;

R6 is H;

R₈ is selected from the group consisting $-C(0)-R_{10}$, $-C(0)-CH_2-OR_{10}, \text{ and } -C(0)CH_2-N(R_{10})(R_{10}), \text{ wherein } R_{10} \text{ is } H, CH_3, \text{ or } -CH_2CH_3;$

 R_{11} is selected from the group consisting of -Ar4, -(CH2) $_{1-3}\text{-Ar}_4,$ and -C(O)-Ar4;

20 R_{15} is -OH or $-OC_{1-4}$ straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, or $-CO_2H$, wherein the R_9 is a $-C_{1-4}$ branched or straight alkyl group, wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O_1 ;

25 Ar₂ is (hh);

Y is O;

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl, thienothienyl, thiadiazolyl, benzotriazolyl, benzo(b)thiophenyl, benzofuranyl, and indolyl, and said

cyclic group optionally being singly or multiply substituted by -Q $_{1}; \\$

each Ar $_4$ cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, naphthyl, pyridinyl, oxazolyl, pyrimidinyl, or indolyl, said cyclic group optionally being singly or multiply substituted by -0_1 ;

each Q_1 is independently selected from the group consisting of -NH2, -Cl, -F, -Br, -OH, -R9, -NH-R5 wherein R_5 is -C(O)-R10 or -S(O)2-R9, -OR5 wherein R_5 is -C(O)-R10, -OR9, -N(R9) (R_{10}) , and

O CH₂,

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Other preferred compounds of embodiment E employ formula (II) wherein R_1 is (el0), X_5 is CH, and the other substituents are as defined above.

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More preferred compounds of embodiment E employ formula (II) wherein R_1 is (e10), X_5 is CH, R_3 is CO-Ar₂, and the other substituents are as defined above.

Other more preferred compounds of embodiment E employ formula (II) wherein R_1 is (e10), X_5 is CH, R_3 is -C(0)-CH₂-T₁-R₁₁, R_{11} is -(CH₂)₁₋₃-Ar₄, and the other substituents are as defined above.

Other more preferred compounds of embodiment E employ formula (II) wherein R_1 is (e10) and X_5 is CH and R_3 is -C(0)-CH₂-T₁-R₁₁, T₁ is 0, R_{11} is -C(0)-Ar₄, and the other substituents are as defined above.

More preferably, in these more preferred compounds, $R_{\tilde{0}}$ is selected from the group consisting of:

15 -C(0)-R₁₀,

5

1.0

-C(0)0-Rq, and

-C(O)-NH-R10.

Alternatively, in these more preferred compounds, R_5 is selected from the group consisting of:

20 -S(0)2-Rg,

-S(O)2-NH-R10,

-C(0)-C(0)-R10,

-Rg, and

-C(0)-C(0)-OR10.

25 Most preferably, in these more preferred compounds,

m 1s 1;

T1 1s 0 or S;

 $\rm R_{15}$ is -OH or -OC $_{1-4}$ straight or branched alkyl

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group optionally substituted with -Ar $_{3,}$ -OH, -OR $_{9}$, or -CO $_{2}$ H, wherein the R $_{9}$ is a -C $_{1-4}$ branched or straight alkyl group, wherein Ar $_{3}$ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O $_{1}$;

5 R₂₁ is -H or -CH₃;

Ar2 is (hh);

Y is O, and

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -01;

each ${\rm Ar_4}$ cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl, said cyclic group being singly or multiply substituted by -0,;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -N(R₉)(R₁₀), and

wherein each R_9 and R_{10} are independently a -C1-6

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straight or branched alkyl group optionally substituted with ${\rm Ar}_3$ wherein ${\rm Ar}_3$ is phenyl;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

Other more preferred compounds of embodiment E employ formula (II) wherein R_1 is (e10), X_5 is CH, R_3 is -C(0)-H, and the other substituents are as defined above.

More preferably, in these more preferred compounds, $R_{\bar{\mathbf{5}}}$ is selected from the group consisting of:

-C(0)-R10,

-C(0)0-Rq, and

-C(O)-NH-R10.

Alternatively, in these more preferred compounds, R_5 is selected from the group consisting of:

-S(0)2-Rq,

-S(O)2-NH-R10,

-C(0)-C(0)-R₁₀,

-Rg, and

-C(0)-C(0)-OR10.

Most preferably, in these more preferred compounds,

25 m is 1;

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 R_{15} is -OH or -OC $_{1-4}$ straight or branched alkyl group optionally substituted with -Ar $_{3}$, -OH, -OR $_{9}$, or -CO $_{2}$ H, wherein the R_{9} is a -CC $_{1-4}$ branched or straight alkyl group, wherein Ar $_{3}$ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O $_{1.7}$

- 144 -

 R_{21} is -H or -CH3;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, said cyclic group optionally being singly or multiply substituted by -01;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -CH, -Rg, -NH-Rg wherein Rg is -C(0)-R₁₀ or -S(0)₂-Rg, -ORg wherein Rg is -C(0)-R₁₀, -ORg, -N(Rg)(R₁₀), and

О /\ СН₂,

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_2 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$,

Other more preferred compounds of embodiment E employ formula (II) wherein R_1 is (e10) and X_5 is CH, R_3 is -CO-CH₂-T₁-R₁₁, and R_{11} is -Ar₄, and the other substituents are as defined above.

More preferably, in these more preferred compounds, $\rm R_{\bar 5}$ is selected from the group consisting of:

- 145 -

-C(O)-NH-R10.

Alternatively, in these more preferred compounds, R₅ is selected from the group consisting of:

$$-S(0)_2-R_9$$
,
 $-S(0)_2-NH-R_{10}$,

 $-c(0)-c(0)-R_{10}$,

 $-R_9$, and

10 $-C(0)-C(0)-OR_{10}$.

Most preferably, in these more preferred compounds,

m is 1;

15

25

 T_1 is 0 or S;

 R_{15} is -OH or a -OC $_{1-4}$ straight or branched alkyl group optionally substituted with -Ar $_3$, -OH, -OR $_9$, or -CO $_2$ H, wherein the R $_9$ is a -C $_{1-4}$ branched or straight alkyl group, wherein Ar $_3$ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

20 R₂₁ is -H or -CH₃;

each Ar₃ cyclic group is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -01;

each Ar4 cyclic group is independently selected

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from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl, said cyclic group optionally being singly or multiply substituted by -01;

each Q₁ is independently selected from the group consisting of $-NH_{2,t}$ -Cl, $-F_t$, -Br, -OH, $-R_9$, $-NH_-R_5$ wherein R_5 is $-C(O) - R_{10}$ or $-S(O) _2 - R_9$, $-OR_5$ wherein R_5 is $-C(O) - R_{10}$, $-OR_4$, $-N(R_9)$ (R_{10}) , and

O CH₂,

1.0

15

20

25

3.0

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar₃ is substituted with a Q_1 group which comprises one or more additional -Ar₃ groups, said additional -Ar₃ groups are not substituted with another -Ar₃.

Other preferred compounds of embodiment E employ formula (II) wherein R_1 is (e10), X_5 is N, and the other substituents are as defined above.

More preferred compounds of embodiment E, employ formula (II) wherein R_1 is (e10), X_5 is N, R_3 is CO-Ar₂, and the other substituents are as defined above.

Other more preferred compounds of embodiment E, employ formula (II) wherein R_1 is (e10), X_5 is N, R_3 is -C(O)-CH₂-T₁-R₁₁, R_{11} is -(CH₂)₁₋₃-Ar₄, and the other substituents are as defined above.

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Other more preferred compounds of embodiment E, employ formula (II) wherein R_1 is (e10) and X_5 is N and:

5
$$R_3$$
 is $-C(0) - CH_2 - T_1 - R_{11}$;
 T_1 is 0; and

 $\ensuremath{\mathtt{R}}_{11}$ is -C(O)-Ar4, and the other substituents are as defined above.

More preferably, in these more preferred compounds, $\rm R_{\bar 5}$ is selected from the group consisting of:

- -C(0)-R₁₀,
 -C(0)O-R₀, and
- -C(0)-NH-R₁₀.

Alternatively, in these more preferred compounds, R₅ is selected from the group consisting of:

- -S(0)2-R9,
- -S(O)2-NH-R10,
- -C(O)-C(O)-R₁₀,
- -Rg, and
- -C(O)-C(O)-OR₁₀.

Most preferably, in these more preferred compounds,

m is 1;

 T_1 is 0 or S;

R₁₅ is -OH or a -OC₁₋₄ straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, or -CO₂H, wherein the R₉ is a -C₁₋₄ branched or straight alkyl group, wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O₁;

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Ar2 is (hh);

Y is O. and

15

20

25

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁:

each Ar₄ cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl, optionally being singly or multiply substituted by -Q;;

each Q_1 is independently selected from the group consisting of -NH $_2$, -Cl, -F, -Br, -OH, -R $_9$, -NH-R $_5$ wherein R $_5$ is -C(0)-R $_{10}$ or -S(0) $_2$ -R $_9$, -OR $_5$ wherein R $_5$ is -C(0)-R $_{10}$, -OR $_9$, -N(R $_9$)(R $_{10}$), and

O / \ CH₂,

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

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Other more preferred compounds of embodiment E, employ formula (II) wherein R_1 is (e10), X_5 is N, R_3 is -C(O)-H, and the other substituents are as defined above.

5 More preferably, in these more preferred compounds, R_5 is selected from the group consisting of:

$$-C(0)O-R_9$$
, and

-C(O)-NH-
$$R_{10}$$
.

10 Alternatively, in these more preferred compounds, R_5 is selected from the group consisting of:

-C(O)-C(O)-OR₁₀.

Most preferably, in these more preferred compounds,

R₁₅ is -OH or -OC₁₋₄ straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, or -CO₂H, wherein the R₉ is a -C₁₋₄ branched or straight alkyl group, wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q₁;

$$R_{21}$$
 is -H or -CH₃;

15

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -01;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀, or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -M(R₉)(R₁₀), and

5

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25

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Other more preferred compounds of embodiment E, employ formula (II) wherein R_1 is (e10), X_5 is N, R_3 is -CO-CH₂-T₁-R₁₁, R_{11} is -Ar₄, and the other substituents are as defined above.

30 More preferably, in these more preferred compounds, R_{\S} is selected from the group consisting of:

$$-C(0)-R_{10}$$
,
 $-C(0)O-R_{9}$, and

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-C(O)-NH-R10.

Alternatively, in these more preferred compounds, R_5 is selected from the group consisting of:

-S(O)2-Rq,

-S(O)2-NH-R10,

-C(O)-C(O)-R₁₀,

 $-R_9$, and

-C(0)-C(0)-OR10.

Most preferably, in these more preferred compounds

10 m is 1;

5

15

T1 is 0 or S;

 R_{15} is -OH or -OC₁₋₄ straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, or -CO₂H, wherein the R_9 is a -C₁₋₄ branched or straight alkyl group, wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

 R_{21} is -H or -CH₃;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl.

benzo{b]thiophenyl, pyridyl, benzofuranyl, and indolyl and said cyclic group optionally being singly or multiply substituted by -Q1;

each Ar₄ cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, or thienyl,

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said cyclic group being singly or multiply substituted by $-Q_1$;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -N(R₉) (R₁₀), and

CH₂

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_4$.

The ICE inhibitors of another embodiment (F) of this invention are those of formula (III):

25 wherein:

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 $\ensuremath{R_{\mathrm{1}}}$ is selected from the group consisting of the following formulae:



- 153 -

(e10)
$$R_{21}$$
 R_{5} R_{5}

5

$$(y2) \qquad \qquad x_7 \xrightarrow{f^2} \qquad \qquad ;$$

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$$\{z\}$$
 $\begin{cases} x_{5} - x_{1} & y_{2} \\ y_{1} & y_{2} \\ y_{3} & y_{4} \end{cases}$; and

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, 5 isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

R₂ is:

10 (b)
$$(\bigcap_{m} OR_{n}$$
 ;

m is 1 or 2;

 $\qquad \qquad \text{each } R_5 \text{ is independently selected from } \\ \text{the group consisting of:} \\$

$$\begin{array}{c} -\text{C(O)} -\text{R}_{10}, \\ -\text{C(O)} -\text{R}_{9}, \\ -\text{C(O)} -\text{N} (\text{R}_{10}) (\text{R}_{10}) \\ -\text{S(O)} _2 -\text{R}_9, \\ -\text{S(O)} _2 -\text{NH} -\text{R}_{10}, \\ -\text{C(O)} -\text{CH}_2 -\text{O} -\text{R}_9, \end{array}$$

-H,

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$$-C(0)C(0)-OR_{10}$$
, and $-C(0)C(0)-N(R_{9})(R_{10})$;

Xs is CH or N;

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$$Y_2$$
 is H_2 or O ;

$$X_7$$
 is $-N(R_8)$ - or -O-;

 $$R_{6}$$ is selected from the group consisting of -H and 10 $$-CH_{3}$;} \label{eq:charge_constraint}$

R₈ is selected from the group consisting of:

-C(0)-R10,

-C(O)O-Rg,

-C(O)-N(H)-R10,

-S(O)2-R9,

-S(O)2-NH-R10,

-C(O)-CH2-OR10,

2 10

-C(0)C(0)-R10;

-C(O)-CH2N(R10)(R10),

-C(O)-CH2C(O)-O-R9,

-C(O)-CH2C(O)-R9,

-H, and

-C(O)-C(O)-OR10;

each R_9 is independently selected from the group consisting of -Ar $_3$ and a - C_{1-6} straight or branched alkyl group optionally substituted with Ar $_3$, wherein the - C_{1-6} alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H., -Ar₃, a C_{3-6} cycloalkyl group, and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is

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optionally unsaturated;

 R_{13} is selected from the group consisting of H, Ar_3 , and a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , -CONH₂, -OR₅, -OH, -OR₆, or -COSH;

each $R_{\rm 21}$ is independently selected from the group consisting of -H or a -C $_{\rm 1-6}$ straight or branched alkyl group;

each R_{51} is independently selected from the group consisting of R_9 , $-C(0)-R_9$, $-C(0)-N(H)-R_9$, or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

each ${\rm Ar_3}$ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, ${\rm SO_2}$, =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

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1.5

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each Q_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, -Q, -OH, -perfluoro C₁₋₃ alkyl, R₅, -OR₅, -NHR₅, OR₉, -N(R₉) (R₁₀), R₉, -C(O)-R₁₀, and O, CH₂,

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provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

Preferred compounds of embodiment F employ formula (III), wherein \mathbf{R}_1 is (w2) and the other substituents are as defined above.

10 Preferably, when R_1 is (w2):

m is 1;

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ring C is benzo, pyrido, or thieno;

 R_5 is selected from the group consisting of:

-C(O)- R_{10} , wherein R_{10} is -Ar₃;

-C(O)O-R₉, wherein R₉ is -CH₂-Ar₃;

-C(O)C(O)- R_{10} , wherein R_{10} is -Ar₃;

 $-R_{\rm 9},$ wherein $R_{\rm 9}$ is a $C_{\rm 1-2}$ alkyl group substituted with $-Ar_{\rm 3};$ and

-C(0)C(0)-OR10, wherein R10 is -CH2Ar3;

R6 is H;

 R_{8} is selected from the group consisting -C(0)-R₁₀, -C(0)-CH₂-OR₁₀, and -C(0)CH₂-N(R₁₀)(R₁₀), wherein R_{1C} is H, CH₃, or -CH₂CH₃;

R₁₃ is H or a C₁₋₄ straight or branched alkyl group optionally substituted with Ar₃, -OH₃, -OO₂H, wherein the R₉ is a C₁₋₄ branched or straight chain alkyl group; wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O₁;

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Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl, thienothienyl, thiadiazolyl, benzotriazolyl, benzo(b)thiophenyl, benzofuranyl, and indolyl.

each Q_1 is independently selected from the group consisting of -NH2, -Cl, -F, -Br, -OH, -R9, -NH-R5 wherein R5 is -C(0)-R10 or -S(0)2-R9, -OR5 wherein R5 is -C(0)-R10, -OR4, -NHR9, and

CH₂,

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wherein each R₉ and R₁₀ are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃ wherein Ar₃ is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_4$.

Other preferred compounds of embodiment F employ formula (III), wherein R_1 is (ell) and the other substituents are as defined above.

Other preferred compounds of embodiment F employ formula (III), wherein ${\bf R}_1$ is (e12) and the other substituents are as defined above.

Other preferred compounds of embodiment F employ formula (III), wherein \mathbf{R}_1 is (y1) and the other substituents are as defined above.

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Other preferred compounds of embodiment F employ formula (III), wherein R_1 is (y2) and the other substituents are as defined above

Other preferred compounds of embodiment F employ formula (III), wherein R_1 is (z) and the other substituents are as defined above.

Other preferred compounds of embodiment F employ formula (III), wherein R_1 is (e10) and X_5 is CH (also referred to herein as e10-B), and the other substituents are as defined above.

Other preferred compounds of embodiment F employ formula (III), wherein R_1 is (e10) and X_5 is N, (also referred to herein as e10-A) and the other substituents are as defined above.

Preferably, when R_1 is (e11), (e12), (y1), (y2), (z), (e10-A), and (e10-B), R_5 is selected from the group consisting of:

```
-C(0)-R-0.
```

Alternatively, when R_1 is (e11), (e12), (y1), (y2), (z), (e10-A), and (e10-B), R_5 is selected from the group consisting of:

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30 More preferably, R₅ is R-C(0)-C(0)-R₁₀.

⁻Rq,

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Alternatively, R_5 is $-C(0)-C(0)-OR_{10}$.

More preferably when R_1 is (e11), (e12), (y1), (y2), (z), (e10-A), and (e10-B):

m is 1:

5 R₂₁ is -H or -CH₃;

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 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar₃, wherein the Ar₃ cyclic group is phenyl, said cyclic group optionally being multiply or singly substituted by $-Q_1$;

10 each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienthienyl, imidazolyl, thiadiazolyl, benzo(b)thiophenyl, pyridyl, benzofuranyl, or indolyl,

benzo[b]thiophenyl, pyridyl, benzofuranyl, or indolyl and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -N(R₉)(R₁₀), and 0 / \(CH₂,

CH

wherein each R_9 and R_{10} are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar_3 , wherein the Ar_3 cyclic group is phenyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

provided that when -Ar $_3$ is substituted with a - \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

More preferably, in these more preferred compounds, the Ar₃ cyclic group is selected from the set consisting of phenyl, naphthyl, thienyl, quinclinyl, isoquinclinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -01.

Compounds in a preferred form of this embodiment F

 R_5 is $-C(0)-R_{10}$, wherein:

 R_{10} is Ar_3 , wherein the Ar_3 cyclic group is phenyl, said cyclic group optionally being singly or multiply substituted by:

-F, -Cl.

5

1.0

15

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2.5

 $-N(H)-R_5$, wherein $-R_5$ is -H or $-C(O)-R_{10}$, wherein

 R_{10} is a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 , wherein the Ar_3 cyclic group is phenyl, said cyclic group optionally being singly or multiply substituted by $-C_1$,

 $-N(R_9)\;(R_{10})\,,$ wherein R_9 and R_{10} are independently a $-\text{C}_{1-4}$ straight or branched alkyl group, or

30 -O-R₅, wherein R₅ is H or a -C₁₋₄ straight or branched alkyl group.

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More preferably the Ar_3 cyclic group is phenyl optionally being singly or multiply substituted at the 3- or 5-position by -Cl or at the 4-position by -NH-Rs, -N(Rg)(Rg)), or -O-Rg.

Other preferred compounds of embodiment F include those wherein R₅ is -C(0)-R₁₀, wherein R₁₀ is Ar₃ and the Ar₃ cyclic group is selected from the group consisting of indolyl, benzimidazolyl, thienyl, and benzo(b)thiophenyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

Other preferred compounds of embodiment F include those wherein R_5 is $-C(0)-R_{10}$, wherein R_{10} is Ar_3 and the Ar_3 cyclic group is selected from quinolyl and isoquinolyl, and said cyclic group optionally being singly or multiply substituted by $-O_1$.

Other preferred compounds of embodiment F are those wherein R_5 is -C(0)- R_{10} , wherein R_{10} is Ar_3 , wherein the Ar_3 cyclic group is phenyl, substituted by

CH₂

In another form of embodiment F the compounds are as described above, further provided that when:

m is 1; R_1 is (e10); X_5 is CH; R_{15} is -OH; R_{2^+} is -H; and

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- $\rm Y_2$ is 0 and $\rm R_3$ is -C(O)-H, then $\rm R_5$ cannot be: -C(O)-R₁₀, wherein $\rm R_{10}$ is -Ar₃ and the Ar₃ cyclic group is phenyl, unsubstituted by -Q₁, 4-(carboxymethoxy)phenyl, 2-fluorophenyl, 2-pyridyl, N-(4-methylpiperazino)methylphenyl, or
- -C(O)-OR $_9$, wherein R $_9$ is -CH $_2$ -Ar $_3$, and the Ar $_3$ cyclic group is phenyl, unsubstituted by -Q $_1$,; and when
- Y_2 is O, R_3 is -C(O)-CH₂-T₁-R₁₁, T₁ is C, and R₁₁ is Ar₄, wherein the Ar₄ cyclic group is 5-(1-(4-chlorophenyl)-3-trifluoromethyl)pyrazolyl), then R_5 cannot be:
- -C(0)- R_{10} , wherein R_{10} is -Ar₃ and the Ar₃ cyclic group is 4-(dimethylaminomethyl)phenyl, phenyl, 4-(carboxymethylthio)phenyl, 4-(carboxymethylthio)phenyl, 4-(carboxypropyl)phenyl, 2-fluorophenyl, 2-pyridyl, N-(4-methylpiperazino)methylphenyl, or
- -C(0)-OR₉, wherein R_9 is -CH₂-Ar₃ and the Ar₃ cyclic group is phenyl;
- and when R_{11} is Ar₄, wherein the Ar₄ cyclic group is 5-(1-phenyl-3-trifluoromethyl)pyrazolyl), then R_5 cannot be:
 - -C(0)-OR9, wherein R9 is -CH2-Ar3, and the Ar3 cyclic group is phenyl;
- and when R_{11} is Ar_4 , wherein the Ar_4 cyclic group is 5-(1-(2-pyridyl)-3-trifluoromethyl)pyrazolyl), then R_5 cannot be:
 - -C(0)-R $_{10}$, wherein R $_{10}$ is -Ar $_{3}$ and the Ar $_{3}$ cyclic group is 4-(dimethylaminomethyl)phenyl, or
 - -C(0)-OR₉, wherein R_9 is -CH₂-Ar₃, and the Ar₂ cyclic group is phenyl, unsubstituted by -Q₁,; and when
 - $\rm Y_2$ is O, R $_3$ is -C(O)-CH $_2$ -T $_1$ -R $_{11}$, T $_1$ is O, and R $_{11}$ is -C(O)-Ar $_4$, wherein the Ar $_4$ cyclic group is 2,5-

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dichlorophenyl, then Rs cannot be:

-C(0)- R_{10} , wherein R_{10} is -Ar₃ and the Ar₃ cyclic group is 4-(dimethylaminomethyl)phenyl, 4-(N-morpholinomethyl)phenyl, 4-(N-

- 5 methylpiperazino)methyl)phenyl, 4 (N-(2-methyl)imidazolylmethyl)phenyl, 5-benzimidazolyl, 5-benztriazolyl, N-carboethoxy-5-benztriazolyl, N-carboethoxy-5-benzimidazolyl, or
- -C(O)-ORg, wherein Rg is -CH2-Ar3, and the Ar3 cyclic group is phenyl, unsubstituted by -Q1,; and when

 Y_2 is H_2 , R_3 is -C(O) $-CH_2-T_1-R_{11}$, T_1 is O, and R_{11} is -C(O) $-Ar_4$, wherein the Ar_4 cyclic group is 2,5-dichlorophenyl, then R_5 cannot be: -C(O) $-OR_9$, wherein R_9 is $-CH_7-Ar_3$ and the Ar_3

In another form of embodiment F, preferred compounds are those wherein R_{21} is -H.

cyclic group is phenyl.

Alternatively, preferred compounds are those wherein $R_{2,1}$ is -CH₃.

Preferred compounds of embodiment F employ formula (III), wherein R_1 is (w2) and the other substituents are as defined above.

More preferably, R1 is (w2) and

25 m is 1:

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ring C is benzo, pyrido, or thieno;

 R_3 is selected from the group consisting of -C(O)-H, -C(O)-Ar_2, and -C(O)CH_2-T_1-R_1;

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 $R_5 \text{ is selected from the group consisting of:} \\ -C(0) -R_{10}, \text{ wherein } R_{10} \text{ is } -Ar_3; \\ -C(0) -R_{10}, \text{ wherein } R_9 \text{ is } -CH_2-Ar_3; \\ -C(0) C(0) -R_{10}, \text{ wherein } R_{10} \text{ is } -Ar_3; \\ -R_9, \text{ wherein } R_9 \text{ is a } C_{1-2} \text{ alkyl group } \\ \text{substituted with } -Ar_3; \text{ and} \\ -C(0) C(0) -OR_{10}, \text{ wherein } R_{10} \text{ is } -CH_2Ar_3; \\ T_1 \text{ is O or S;} \\ \\ 10 \qquad \qquad R_6 \text{ is } H; \\ \\ \end{array}$

R₆ is H;

20

 R_8 is selected from the group consisting -C(0)-R₁₀, -C(0)-CH₂-OR₁₀, and -C(0)CH₂-N(R₁₀) (R₁₀), wherein R₁₀ is H, CH₃, or -CH₂CH₃;

 $$\rm R_{11}$ is selected from the group consisting of -Ar4,

 R_{15} is -OH or $-OC_{1-4}$ straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, or -CO₂H, wherein the R_9 is a -Cl₁₋₄ branched or straight alkyl group, wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with $Q_{1,7}$

 Ar_2 is (hh);

Y is 0;

each Ar₃ cyclic group is independently selected
from the set consisting of phenyl, naphthyl, thienyl,
quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl,
thienothienyl, thiadiazolyl, benzotriazolyl,
benzo[b]thiophenyl, benzofuranyl, and indolyl, and said
cyclic group optionally being singly or multiply
substituted by -Q₁;

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each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, naphthyl, pyridinyl, oxazolyl, pyrimidinyl, or indolyl, said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -N(R₉) (R₁₀), and

CH₂,

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wherein each R₉ and R₁₀ are independently a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃ wherein Ar₃ is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Other preferred compounds of embodiment F employ formula (III), wherein \mathbf{R}_1 is (ell) and the other substituents are as defined above.

Other preferred compounds of embodiment F employ formula (III), wherein R_1 is (e12) and the other substituents are as defined above

Other preferred compounds of embodiment F employ formula (III) wherein R_1 is (y1) and the other substituents are as defined above.

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Other preferred compounds of embodiment F employ formula (III) wherein \mathbf{R}_1 is (y2) and the other substituents are as defined above.

Other preferred compounds of embodiment F of 5 employ formula (III) wherein R₁ is (z) and the other substituents are as defined above.

Other preferred compounds of embodiment F employ formula (III) wherein R_1 is (e10), X_5 is CH, and the other substituents are as defined above.

Other preferred compounds of embodiment F employ formula (III) wherein R_1 is (e10), X_5 is N, and the other substituents are as defined above.

More preferably, in these more preferred compounds, ${\rm R}_{\bar 5}$ is selected from the group consisting of:

- 15 -C(O)-R₁₀,
 - -C(0)0-Rq, and
 - -C(O)-NH-R10.

Alternatively, in these more preferred compounds, R₅ is selected from the group consisting of:

20 -S(0)₂-R₉,

25

- -S(O)2-NH-R10,
- -C(0)-C(0)-R10,
- -Ra,
- $-C(0)-C(0)-OR_{10}$, and
- -C(O)C(O)-N(R9)(R10).

Most preferably, in these more preferred compounds,

m is 1:

 R_{13} is H or a $-C_{1-4}$ straight or branched alkyl

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group optionally substituted with -Ar $_3$, -OH, -OR $_9$, or -CO $_2$ H, wherein the R $_9$ is a -C $_{1-4}$ branched or straight alkyl group, wherein Ar $_3$ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O_{1} ;

5 R₂₁ is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl, optionally substituted by -Q-;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -N(R₉)(R₁₀), and

20

25

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wherein each R_9 and R_{10} are independently a $-C_{1-\delta}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted

- 169 -

with another -Ar3.

 $\label{eq:preferred compounds of embodiment (F)} include, but are not limited to:$

2001

5 2100a

2100b

2100c

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2100d

2100e

The ICE inhibitors of another embodiment (G) of this invention are those of formula (IV):

(IV) (IV) (IV) (R₁-N R₃

wherein:

m is 1 or 2;

10 $$R_{1}$$ is selected from the group consisting of the following formulae:

(ell)
$$\begin{array}{c} Y_2 \\ \\ R_5 - N \\ H \end{array}$$

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(z)
$$\begin{array}{c} x_{1} \\ x_{2} \\ R_{5} - N \\ H \end{array} \hspace{0.5cm} \text{; and}$$

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

R₃ is selected from the group consisting of: -CN, -C(0)-H, -C(0)-CH₂-T₁-R₁₁, -C(0)-CH₂-F, -C=N-O-R₉, and -CO-Ar₂;

each R_5 is independently selected from the $\,$

15 group consisting of:
$$-C(0) - R_{10}, \\ -C(0) - R_{10}, \\ -C(0) - R_{10}, \\ -C(0) - R_{10}, \\ (R_{10}) (R_{10}) \\ -S(0) _{2} - R_{9}, \\ -S(0) _{2} - NH - R_{10}, \\ -C(0) - CH_{2} - O - R_{9}, \\ -C(0) C(0) - R_{10}, \\ -R_{9}, \\ -H, \\ 25 \\ -C(0) C(0) - N(R_{9}) (R_{10}), \\ and \\ -C(0) C(0) - N(R_{9}) (R_{10}), \\ (R_{10}), \\ (R_{10}) (R_{10}) (R_{10}) (R_{10}), \\ (R_{10}) (R_{10}) (R_{10}) (R_{10}) (R_{10}) (R_{10}) (R_{10}) (R_{10}), \\ (R_{10}) (R_{10}$$

 Y_2 is H_2 or O;

$$X_7$$
 is $-N(R_8)$ - or -O-;

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each T_1 is independently selected from the group consisting of -O-, -S-, -S(O)-, and -S(O)2-;

 $$R_{6}$$ is selected from the group consisting of -H and 5 -CHa;

 $\ensuremath{\text{R}_8}$ is selected from the group consisting of:

-C(O)-R10,

-C(0)0-Rg,

10 -C(0)-NH-R₁₀,

15

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25

-S(0)2-R9,

-S(O)2-NH-R10,

-C(O)-CH2-OR10,

-C(0)C(0)-R10,

 $C(0)C(0)-R_{10}$

 $-C(0)-CH_2-N(R_{10})(R_{10})$,

-C(O)-CH2C(O)-O-R9,

-C(O)-CH₂C(O)-R₉,

-H, and

-C(O)-C(O)-OR10;

each R_9 is independently selected from the group consisting of -Ar₃ and a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar₃, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

 $\qquad \qquad \text{each R_{11} is independently selected from the group} \\ \text{30} \qquad \text{consisting of:} \\$

-Ara,

-(CH₂)₁₋₃-Ar₄,

-H, and

- 174 -

-C(0)-Ar4;

5

1.0

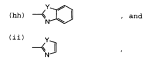
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 R_{15} is selected from the group consisting of -OH, -OAr₃, -N(H)-OH, and -OC₁₋₆, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with Ar_3 , -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

each $\rm R_{21}$ is independently selected from the group consisting of -H or a -C $_{1-6}$ straight or branched alkyl group;

 \mbox{Ar}_2 is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $\mbox{-Q}_1$ or phenyl, optionally substituted by \mbox{Q}_1 :



wherein each Y is independently selected from the group consisting of O and S;

each Ar $_3$ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO $_2$, =N-, and -NH-, -N(R $_3$)-, and -N(R $_9$)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or

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multiply substituted by -Q1;

each Ar_4 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q_1 is independently selected from the group consisting of -NH2, -CO_2H, -Cl, -F, -Br, -I, -NO_2, -CN, =O, -OH, -perfluoro C_{1-3} alkyl, R_5, -OR_5, -NHR_5, OR_9, -N(R_9) (R_{10}), R_9, -C(O)-R_{10}, and O CH_2;

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1.5

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provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$;

Preferred compounds of embodiment G employ formula (IV), wherein R_1 is (w2) and the other substituents are as defined above.

Preferably, when R1 is (w2):

30 m is 1;

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ring C is benzo, pyrido, or thieno:

 R_5 is selected from the group consisting of:

-C(O)-R₁₀, wherein R₁₀ is -Ar₃;

-C(O)O-R₉, wherein R₉ is -CH₂-Ar₃;

-C(0)C(0)-R₁₀, wherein R₁₀ is -Ar₃;

 $-R_9$, wherein R_9 is a C_{1-2} alkyl group

substituted with -Ar3; and

-C(O)C(O)-OR₁₀, wherein R₁₀ is -CH₂Ar₃;

10 R₆ is H;

5

15

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 R_8 is selected from the group consisting -C(0)-R₁₀, -C(0)-CH₂-OR₁,, and -C(0)CH₂-N(R₁₀)(R₁₀), wherein R₁₀ is H, CH₃, or -CH₂CH₃;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with Ar₃, -OH, -OR₉, -CO₂H, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

Ar₃ is phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl, thienothienyl, thiadiazolyl, benzotriazolyl, benzo[b]thiophenyl, benzofuranyl, and indolyl;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₀, and

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

5 provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Other preferred compounds of embodiment G employ formula (IV) wherein R_1 is (e10-A) and the other substituents are as defined above.

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Other preferred compounds of embodiment G employ formula (IV) wherein R_1 is (ell) and the other substituents are as defined above.

Other preferred compounds of embodiment G employ formula (IV) wherein R_1 is (e12) and the other substituents are as defined above.

Other preferred compounds of embodiment G employ formula (IV) wherein R_1 is (y1) and the other substituents are as defined above.

Other preferred compounds of embodiment G employ formula (IV) wherein R_1 is (y2) and the other substituents are as defined above.

Other preferred compounds of embodiment G employ formula (IV) wherein R_1 is (z) and the other substituents are as defined above.

More preferred compounds of embodiment G are those wherein $R_{\rm 3}$ is -CO-Ar₂.

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Most preferably, when R₃ is -CO-Ar₂, Y is O.

Other more preferred compounds are those wherein R_3 is -C(0) $-CH_2-T_1-R_{11}$ and R_{11} is $-(CH_2)_{1-3}-Ar_4$.

Most preferably, when R_3 is -C(0)-CH₂-T₁-R₁₁ and R_{11} is -(CH₂)₁₋₃-Ar₄, T_1 is 0.

Other more preferred compounds are those wherein: $\begin{array}{l} R_3 \text{ is } -\text{C}(0) - \text{CH}_2 - \text{T}_1 - R_{11}; \\ T_1 \text{ is 0; and} \\ R_{11} \text{ is } -\text{C}(0) - \text{Ar}_4 \,. \end{array}$

Other more preferred compounds are those wherein $\rm R_3$ is -C(O)-H.

Other more preferred compounds are those wherein R_3 is -CO-CH $_2$ -T $_1$ -R $_{11}$ and R_{11} is -Ar $_4$.

More preferably, when $\rm R_3$ is -CO-CH $_2$ -T $_1$ -R $_{11}$ and 15 $\rm R_{11}$ is -Ar $_4$, T $_1$ is 0 or S.

More preferably, when $R_1,$ is (ell), (el2), (y1), (y2), (z), (el0-A), and (el0-B), R_5 is selected from the group consisting of:

-C(O)-R-0,

-C(0)0-R9, and

-C(O)-NH-R10.

Alternatively, when R_1 , is (e11), (e12), (y1), (y2), (z), (e10-A), and (e10-B), R_5 is selected from the group consisting of:

-S(O)₂-R₉,

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-S(O)2-NH-R10,

-C(O)-C(O)-R10,

-Ra,

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-C(0)-C(0)-OR10, and -C(0)-C(0)-N(R9)(R10).

More preferably, Rs is -C(0)-C(0)-R10.

Alternatively, R5 is -C(0)-C(0)-OR10.

Most preferably, when R_1 is (e11), (e12), (y1), (y2), (z), (e10-A), and (e10-B),;

m is 1:

 R_{21} is -H or -CH₃;

R51 is a C1-6 straight or branched alkyl group optionally substituted with Ar3, wherein the Ar3 cyclic group is phenyl, said cyclic group optionally being multiply or singly substituted by -Q1;

each Ara cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, or indolyl, and said cyclic group optionally being singly or multiply substituted by -Q1;

each Q1 is independently selected from the group consisting of -NH2, -Cl, -F, -Br, -OH, -Rg, -NH-Rg wherein R_5 is $-C(0)-R_{10}$ or $-S(0)_2-R_9$, $-OR_5$ wherein R_5 is $-C(0)-R_{10}$, $-OR_9$, $-N(R_9)(R_{10})$, and

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1.5

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 , wherein the Ar_3 cyclic group is phenyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

provided that when -Ar $_3$ is substituted with a -Q $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_4$.

More preferably, in these more preferred compounds, the λr_3 cyclic group is selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo [b] thiophenyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.

Compounds in a preferred form of embodiment G are those wherein R_{21} is H and the other substituents are as defined above.

Compounds in another preferred form of embodiment G are those wherein R_{21} is CH_3 and the other substituents are as defined above.

The ICE inhibitors of another embodiment (H) of this invention are those of formula (V):

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$$(V)$$
 (I) R_1 R_2 R_3

wherein:

m is 1 or 2;

5 R₁ is:

R3 is selected from the group consisting of:

-CN,

-CO-Ar2;

each R_5 is independently selected from the group

15 consisting of:

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$$-C(0)-R_{10}$$
,

$$-\text{C(O)C(O)}-\text{N(R}_9)\text{ }(\text{R}_{10})\text{ , and}$$

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 Y_2 is H_2 or O;

-H, and -C(0)-C(0)-OR₁₀;

each T_1 is independently selected from the group consisting of -O-, -S-, -S(O)-, and -S(O)₂-;

each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a - C_{1-6} straight or branched alkyl group optionally substituted with Ar₃, wherein the - C_{1-6} alkyl group is optionally unsaturated:

each $\ensuremath{R_{\text{I}1}}$ is independently selected from the group consisting of:

-Ar₄,
-(CH₂)₁₋₃-Ar₄,
-H, and
-C(O)-Ar₄;

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- 183 -

 R_{15} is selected from the group consisting cf -OH, -OAr₃, -N(H)-OH, and -OC₁₋₆, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with Ar₁,

-CONH2, -OR5, -OH, -OR9, or -CO2H;

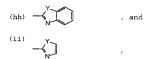
R₂₁ is -CH₃;

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Ar₂ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by Q_1 :



wherein each Y is independently selected from the \$15\$ group consisting of O and $S_{\it f}$

each Ar₃ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

each Ar4 is a cyclic group independently selected

from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -0-, -S-, -SO-, SO₂, =N-, -NH-, -N(\mathbb{R}_5)-, and -N(\mathbb{R}_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$;

Compounds of another form of embodiment (I) (form 1) are those of formula (V):

$$(V)$$
 $(P)_{m} R_{5}$ R_{1} R_{3}

wherein.

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m is 1 or 2;

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$$R_3 \text{ is selected from the group consisting of:} \\ -CN, \\ -C(O) - H, \\ -C(O) - CH_2 - T_1 - R_{11}, \\ -C(O) - CH_2 - F, \\ -C - N - O - R_9, \text{ and } \\ -CO - AT_2; \\ \\ 10 \text{ each } R_5 \text{ is } -C(O)C(O) - OR_{10}; \\ Y_2 \text{ is } H_2 \text{ or } O; \\ \text{ each } T_1 \text{ is independently selected from the group consisting of } -O -, -S -, -S(O) -, \text{ and } -S(O)_2 -; \\ \\ 15 \text{ } R_8 \text{ is selected from the group consisting of:} \\ -C(O) - R_{10}, \\ -C(O) - NH - R_{10}, \\ -C(O) - NH - R_{10}, \\ -S(O)_2 - R_9, \\ -S(O)_2 - R_9, \\ -S(O)_2 - NH - R_{10}, \\ -C(O) - CH_2 - OR_{10}, \\ -C(O) - CH_2 - CO) - OR_9, \\ -C(O) - CH_2 - CO) - CR_9, \\ -C(O) - CR_2 - CO) - CR_9, \\ -C(O) - CR_9 - CCO) -$$

-H, and -C(O)-C(O)-OR₁₀;

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each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

5 each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar₃, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

10 each R_{11} is independently selected from the group consisting of:

-Ar₄,
- (CH₂)₁₋₃-Ar₄,
-H, and
-C(0)-Ar₄;

1.5

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 R_{15} is selected from the group consisting of -OH, -OAr_3, -N(H)-OH, and -OC_{1-6}, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with Ar_3, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

each R₂₁ is independently selected from the group consisting of -H or a -C₁₋₆ straight or branched alkyl group;

Ar₂ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by Q_1 :

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wherein each Y is independently selected from the
group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O1;

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each Ar_4 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

30 each Q₁ is independently selected from the group

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consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =0, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, OR_9 , -N(R_9)(R_{10}), R_9 , -C(O)- R_{10} , and O CH₂;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_2$:

Alternatively, compounds of this form of embodiment I (form 2) are those wherein $R_{2,1}$ is -CH₃.

Compounds of another form of embodiment (J) (form 15 $\,$ 1) are those of formula (V):

wherein:

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m is 1 or 2;

20 R₁ is:

 R_3 is selected from the group consisting of: -CN, -C(O)-H,

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-C(0)-CH2-T1-R11,
                   -C(O)-CH2-F,
                   -C=N-O-R_9, and
                   -CO-Ar2;
 5
                  each R_5 is independently selected from the
       group consisting of:
                   -C(O)-R10,
                  -C(0)0-Rq,
                  -C(0)-N(R10)(R10)
10
                  -S(O)2-Rq,
                  -S(0)_2-NH-R_{10},
                  -C(O)-CH2-O-R9,
                  -C(0)C(0)-R<sub>10</sub>,
                  -Rq,
15
                  -H,
                  -C(0)C(0)-OR10, and
                  -C(0)C(0)-N(R9)(R10);
            Yo is Ho or O;
            each T_1 is independently selected from the group
      consisting of -O-, -S-, -S(O)-, and -S(O)2-;
20
            R_8 is selected from the group consisting of:
                  -C(O)-R10.
                  -C(O)O-Rq,
25
                  -C(O)-NH-R10,
                  -S(O)2-Rg,
                  -s(0)_2-NH-R_{10},
                  -C(O)-CH2-OR10,
                  -C(0)C(0)-R10,
                 -C(0) - CH_2 - N(R_{10})(R_{10}),
30
                  -C(O)-CH-C(O)-O-Rq,
                  -C(0)-CH-C(0)-Rq,
                  -H.
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 $-C(0) - C(0) - OR_{10}$, and $-C(0) - C(0) - N(R_9)(R_{10})$;

each R_9 is independently selected from the group, consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 , wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

each \mathbf{R}_{11} is independently selected from the group consisting of:

-Ara,

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-(CH2)1-3-Ar4,

-H, and

-C(0)-Ara:

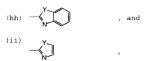
 R_{15} is selected from the group consisting of -OH, -OAr3, -N(H)-OH, and -OC1-6, wherein C1-6 is a straight or branched alkyl group optionally substituted with Ar3,

-CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

each R_{21} is independently selected from the group consisting of -H or a $-C_{1-6}$ straight or branched alkyl group;

 $\rm Ar_2$ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $\rm -Q_1$ or phenyl, optionally substituted by $\rm Q_1$:





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wherein each Y is independently selected from the group consisting of O and S:

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O1;

each Ar_4 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O1;

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consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =0, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NH R_5 , OR₉, -N(R_9) (R_{10}), R_9 , -C(0)-R₁₀, and O CH₂;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$;

provided that when:

m is 1;
R₁ is (e10);
15 X₅ is CH;
R₁₅ is -OH;
R₂₁ is -H; and

cannot be:

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 Y_2 is 0, R_3 is -C(0)-CH₂-T₁-R₁₁, T_1 is 0, and R_{11} is A_{14} , wherein the A_{14} cyclic group is 5-(1-(4-chlorophenyl)-3-trifluoromethyl)pyrazolyl), then R_5

-C(0)-R₁₀, wherein R₁₀ is -Ar₃ and the Ar₃ cyclic group is 4-(dimethylaminomethyl)phenyl, phenyl, 4-(carboxymethylthio)phenyl,4-(carboxyethylthio)phenyl, 4-(carboxyethyl)phenyl, 4-(carboxypropyl)phenyl, 2-

- 193 -

-C(O)-ORq, wherein Rq is -CH2-Ar3 and the Ar3

and when R_{11} is Ar_4 , wherein the Ar_4 cyclic group is 5-(1-phenyl-3-trifluoromethyl)pyrazolyl), then R_5

fluorophenyl, 2-pyridyl, N-(4-methylpiperazino)methylphenyl, or

cyclic group is phenyl:

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cannot be:
     -C(O)-ORq, wherein Rq is -CH2-Ar3, and the Ar3
cyclic group is phenyl;
     and when R11 is Ar4, wherein the Ar4 cyclic group
is 5-(1-(2-pyridyl)-3-trifluoromethyl)pyrazolyl), then
Rs cannot be:
     -C(O)-R10, wherein R10 is -Ar3 and the Ar3 cyclic
group is 4-(dimethylaminomethyl)phenyl, or
     -C(O)-OR9, wherein R9 is -CH2-Ar3, and the Ar3
cyclic group is phenyl, unsubstituted by -Q1,; and when
     Y_2 is 0, R_3 is -C(0)-CH_2-T_1-R_{11}, T_1 is 0, and R_{11}
is -C(O)-Ar4, wherein the Ar4 cyclic group is 2,5-
dichlorophenyl, then Rs cannot be:
     -C(O)-R10, wherein R10 is -Ar2 and the Ar2 cyclic
group is 4-(dimethylaminomethyl)phenyl, 4-(N-
morpholinomethyl) phenyl, 4-(N-
methylpiperazino) methyl) phenyl, 4-(N-(2-
methyl)imidazolylmethyl)phenyl, 5-benzimidazolyl, 5-
benztriazolyl, N-carboethoxy-5-benztriazolyl, N-
carboethoxy-5-benzimidazolyl, or
     -C(O)-ORq, wherein Rq is -CH2-Arg, and the Arg
cyclic group is phenyl, unsubstituted by -Q1,; and when
     Y_2 is H_2, R_3 is -C(0)-CH_2-T_1-R_{11}, T_1 is O, and R_{11}
is
-C(O)-Ar4, wherein the Ar4 cyclic group is 2,5-
dichlorophenyl, then Rs cannot be:
     -C(O)-OR9, wherein R9 is -CH2-Ar3 and the Ar3
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cyclic group is phenyl.

Compounds of another form of embodiment J (form 2) are those wherein $R_{\rm 21}$ is -CH3.

Compounds of another form of embodiment J (form 3) 5 are those wherein R_5 is -C(O)-C(O)-OR₁₀.

Compounds of another form of embodiment J (form 4) are those wherein R_5 is $-C(0)-C(0)-OR_{10}$ and R_{21} is $-CH_3$.

Preferred compounds of embodiments H, I, and J employ formula (V), wherein R $_3$ is -CO-Ar $_2$.

More preferably, when R₃ is -CO-Ar₂ Y is O.

Preferred compounds of embodiments H, I, and J employ formula (V), wherein R₃ is $-C(O)-CH_2-T_1-R_{11}$ and R₁₁ is $-(CH_2)_{1-3}-Ar_4$.

More preferably, when R $_3$ is -C(0)-CH $_2$ -T $_1$ -R $_{11}$ and R $_{11}$ is -(CH $_2$) $_1$ -3-Ar $_4$, T $_1$ is 0.

Preferred compounds of embodiments H, I, and J employ formula (V), wherein R $_3$ is -C(O)-CH $_2$ -T $_1$ -R $_{11}$, T $_1$ is O, and R $_{11}$ is -C(O)-Ar $_4$.

Preferred compounds of embodiments H, I, and J employ formula (V), wherein R $_3$ is -C(O)-H.

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Preferred compounds of embodiments H, I, and J employ formula (V), wherein R_3 is -CO-CH_2-T_1-R_{11} and R_{11} is -Ar_4.

More preferably, when R_3 is -CO-CH2-T1-R11 and

- 195 -

 \mathtt{R}_{11} is -Ar4, \mathtt{T}_1 is 0 or S.

More preferred compounds of embodiments H and J (forms 1 and 2) are those wherein $R_{\bar{\bf 5}}$ is selected from the group consisting of:

-C(O)-R₁₀,

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-C(0)0-Ra, and

-C(0)-NH-R₁₀.

Alternatively, more preferred compounds of embodiments H and J (forms 1 and 2) are those wherein R_5 is selected from the group consisting of:

-S(O)2-R9,

 $-s(0)_2-NH-R_{10}$,

-C(0)-C(0)-R₁₀,

-R₉,

 $-C(0)-C(0)-OR_{10}$, and

-C(O)-C(O)-N(R₀)(R₁₀).

Most preferably, R_5 is $-C(0)-C(0)-R_{10}$.

Alternatively, R_5 is $-C(0)-C(0)-OR_{10}$.

More preferred compounds of embodiments H, I (form 2), and J (forms 2 and 4) are those wherein:

m is 1;

Y2 is 0;

 R_{15} is -OH or -OC $_{1-4}$ straight or branched alkyl group optionally substituted with Ar $_3$, -OH, -OR $_9$, -CO $_2$ H, wherein the R $_9$ is a C $_{1-4}$ branched or straight chain alkyl group; wherein Ar $_3$ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O $_1$;

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Ar2 is (hh);

Y is O, and

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinclinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Ar₄ cyclic group is independently selected from the group consisting of phenyl, tetrazolyl, pyridyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q1 is independently selected from the group consisting of -NH2, -C1, -F, -Br, -OH, -R9, -NH-R5 wherein R5 is -C(0)-R10 or -S(0)2-R9, -OR5 wherein R5 is -C(0)-R10, -OR9, -N(R9)(R10), and 0 /\ CH2,

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3.0

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein the Ar_3 cyclic group is phenyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

provided that when -Ar $_3$ is substituted with a \mathcal{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted

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with another -Ar3.

More preferred compounds of embodiments I (form 1), and J (form 3) are those wherein:

m is 1;

5 R₂₁ is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar₃, wherein the Ar₃ cyclic group is phenyl, said cyclic group optionally being multiply or singly substituted by $-Q_1$;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, or indolyl, and said cyclic group optionally being singly or multiply substituted by -01;

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3.0

20

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 , wherein the Ar_3 cyclic group is phenyl, and said cyclic group optionally being singly or multiply substituted by $-O_{11}$;

- 198 -

provided that when -Ar $_3$ is substituted with a -Q $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Preferably, in these more preferred compounds the Ar_3 cyclic group is selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo [b]thiophenyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.

Preferred compounds of embodiments H, and J (forms 1 and 1) are those wherein:

 R_3 is $-C(0)-CH_2-T_1-R_{11}$; T_1 is 0; and

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 R_{11} is -C(0)-Ar4, wherein the Ar4 cyclic group is selected from the set consisting of tetrazolyl, pyridyl, oxazolyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by -Q₁.

Preferred compounds of embodiments H, I, and J employ formula (V), wherein R_3 is -CO-CH₂-T₁-R₁₁, R_{11} is -Ar₄, wherein the Ar₄ cyclic group is pyridyl, and said cyclic group optionally being singly or multiply substituted by -Q₁.

Preferred compounds of embodiment J (form 1) are those wherein:

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 R_5 is -C(O)- R_{10} , wherein:

 R_{10} is Ar_3 , wherein the Ar_3 cyclic group is phenyl optionally being singly or multiply substituted by:

-F.

-Cl.

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1.5

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 $-N(H)-R_5$, wherein $-R_5$ is -H or $-C(0)-R_{10}$, wherein R_{10} is a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 , wherein Ar_3 is phenyl,

 $^{-N}(R_9)\,(R_{10})\,,$ wherein R_9 and R_{10} are independently a $^{-C_{1-4}}$ straight or branched alkyl group, or

 $-O-R_5$, wherein R_5 is H or a $-C_{1-4}$ straight or branched alkyl group.

More preferably, Ar_3 is phenyl being optionally singly or multiply substituted at the 3- or 5-position by -Cl or at the 4-position by -NH-R₅, -N(R₉) (R₁₀), or -O-R₆.

Other more preferred compounds of embodiment J (form 1) are those wherein:

 R_3 is -C(0)-H;

 R_5 is -C(0)- R_{10} , wherein R_{10} is Ar_3 and the Ar_3 cyclic group is selected from the group consisting of is indolyl, benzimidazolyl, thienyl, and benzo[b]thiophenyl, and said cyclic group optionally being singly or multiply substituted by - Q_1 ;

Other more preferred compounds of embodiment J (form 1) are those wherein:

R₃ is -C(0)-H;

 R_5 is -C(0)- R_{10} , wherein R_{10} is Ar_3 and the Ar_3

- 200 -

cyclic group is selected from quinolyl and isoquinolyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.

Other more preferred compounds of embodiment J (form 1) are those wherein:

 R_3 is -C(0)-H;

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 $R_{\rm 5}$ is -C(O)-R₁₀, wherein $R_{\rm 10}$ is $Ar_{\rm 3}$ and the $Ar_{\rm 3}$ cyclic group is phenyl, substituted by

O CH2

Preferred compounds of embodiment (J) include, but are not limited to:

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2201

The ICE inhibitors of another embodiment (K) of this invention are those of formula:

wherein:

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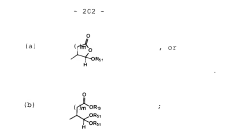
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R₁ is:

(e10)
$$\begin{array}{c} R_{21} \\ R_{5} - N \\ R_{5} - N \end{array}$$
, or
$$\begin{pmatrix} R_{6} \\ R_{5} - N \\ R_{5$$

C is a ring chosen from the set consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl; the ring optionally being singly or multiply substituted by $-Q_1$;

R₂ is:



m is 1 or 2;

5 each $R_{\bar{\mathbf{5}}}$ is independently selected from the group consisting of:

$$\begin{array}{c} -C(O) - R_{10}, \\ -C(O) - R_{9}, \\ -C(O) - N(R_{10}) (R_{10}) \\ 10 & -S(O)_2 - R_9, \\ -S(O)_2 - NH - R_{10}, \\ -C(O) - CH_2 - O - R_9, \\ -C(O) C(O) - R_{10}, \\ -R_9, \\ -H, \\ -C(O) C(O) - OR_{10}, \text{ and } \\ -C(O) C(O) - N(R_9) (R_{10}), \end{array}$$

X5 is CH or N;

$$Y_2$$
 is H_2 or O ;

 $R_{\tilde{\mathbf{G}}}$ is selected from the group consisting of -H and -CH $_{3};$

25 R₈ is selected from the group consisting of:

- 203 -

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each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of $^{-}$ H, $^{-}$ Ar $_3$, a $^{-}$ C $_3$ - $_6$ cycloalkyl group, and a $^{-}$ Cl $_{-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein the $^{-}$ Cl $_{-6}$ alkyl group is optionally unsaturated;

 R_{13} is selected from the group consisting of H, Ar $_3$, and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, -CONH $_2$, -OR $_5$, -OH $_6$, -OR $_9$, or -CO $_2$ H;

each R_{51} is independently selected from the group consisting of R_9 , $-C(0)-R_9$, $-C(0)-N(H)-R_9$, or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

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each R_{21} is independently selected from the group consisting of -H or a - C_{1-6} straight or branched alkyl group;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q₁ is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =O, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, -OR₉, -N(R₉) (R₁₀), -R₉, -C(O)-R₁₀, and O

 $N(R_9)$ (R_{10}) , $-R_9$, -C(0) $-R_{10}$, and O / CH_2 ,

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

30 Preferred compounds of this embodiment are those wherein:

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- 205 -

C is a ring chosen from the set consisting of benzo, pyrido, or thieno the ring optionally being singly or multiply substituted by halogen, -NH $_2$, -NH-R $_3$, -OR $_{10}$, or -R $_9$, wherein R $_9$ is a straight or branched C $_{1-4}$ alkyl group and R $_{10}$ is H or a straight or branched C $_{1-4}$ alkyl group;

Re is H;

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 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with Ar_{2} , -OH, $-OR_{9}$, $-CO_{2}H$, wherein the R_{9} is a C_{1-4} branched or straight chain alkyl group; wherein Ar_{3} is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O_{1} ;

 R_{21} is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with Ar₃, wherein Ar₃ is phenyl, optionally substituted by $-Q_1$;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl benzefuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NiR₉, and

- 206 -

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

 $\hbox{ Preferably, in this preferred embodiment, R_1 is (w2) and the other substituents are as defined above. }$

Compounds of this preferred embodiment include, but are not limited to:

More preferably, R_8 is selected from the

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1.0

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group consisting of:
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-C(0)-R10,

-C(0)0-Rg,

-C(0)-CH $_2$ -OR $_{10}$, and

-C(O)-CH2C(O)-R9.

Most preferably, R_8 is $-C(0)-CH_2-$

 $\ensuremath{\text{OR}_{10}}$ and $\ensuremath{\text{R}_{10}}$ is -H or -CH3.

Alternatively, in this preferred embodiment, R_1 is (el0) and X_5 is CH and the other substituents are as defined above.

Alternatively, in this preferred embodiment, ${\bf R}_1$ is (e10) and ${\bf X}_5$ is N and the other substituents are as defined above.

Preferably, in any of the above compounds of the embodiment (K), R_5 is $-C(0)-R_{10}$ or $-C(0)-C(0)-R_{10}$ and the other substituents are as defined above.

More preferably, $\ensuremath{\mathrm{R}}_{10}$ is $-\ensuremath{\mathrm{Ar}}_3$ and the other substituents are as defined above.

More preferably, in these more preferred

20 compounds:

 R_5 is $-C(0)-R_{10}$ and R_{10} is Ar_{30}

wherein the Ar₃ cyclic group is phenyl optionally being singly or multiply substituted by:

 $-R_{\rm 9},$ wherein $R_{\rm 9}$ is a $C_{\rm 1-4}$ straight or branched 25 $\,$ alkyl group;

-F.

-Cl,

 $^{-N}(H)\!-\!R_5$, wherein $^{-}\!R_5$ is $^{-}\!H$ or $^{-}\!C\,(O)\!-\!R_{10}$, wherein R_{10} is a $^{-}\!C_{1-6}$ straight or branched alkyl group

30 optionally substituted with Ar₃, wherein Ar₃ is phenyl,

 $^{-N}\left(R_{9}\right)\left(R_{10}\right),$ wherein R_{9} and R_{10} are independently a $^{-C_{1-d}}$ straight or branched alkyl group, or

-O-R5, wherein R_5 is H or a -C $_{1-4}$ straight or branched alkyl group.

- 208 -

Preferred compounds of this more preferred embodiment include, but are not limited to:

- 209 -

Most preferably, Ar $_3$ is phenyl being singly or multiply substituted at the 3- or 5-position by -Cl or at the 4-position by -NH-R $_5$, -N(R $_9$) (R $_{10}$), or -O-R $_5$.

Preferred compounds of this most preferred embodiment include, but are not limited to:

- 210 -

Other preferred compounds of this most preferred embodiment include, but are not 5 limited to:

Alternatively, Ar_3 is phenyl being singly or multiply substituted at the 3- or 5-position by $-R_9$, wherein R_9 is a C_{1-4} straight or branched alkyl group; and at the 4-position by $-0-R_5$.

Preferred compounds of this most preferred embodiment include, but are not limited to:

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- 212 -

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 $\\ \qquad \qquad \text{Other preferred compounds of } \\ \text{this most preferred embodiment include, but are } \\ \text{nct} \\$

limited to:

- 214 -

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Alternatively, in this more preferred embodiment, R_5 is $-C(0)-R_{10}$, wherein R_{10} is Ar_3 and the Ar_3 cyclic group is selected from the group consisting of is indolyl, benzimidazolyl, thienyl, quinolyl, isoquinolyl and benzo[b]thiophenyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.

 $\label{eq:most_preferably} \mbox{Most preferably, the Ar_3 cyclic group is isoquinoly1.}$

Preferred compounds of this most preferred embodiment include, but are not limited to:

- 215 -

- 216 -

Other preferred compounds of this most preferred embodiment include, but are not limited to:

- 217 -

- 219 -

Alternatively, in this more preferred embodiment, R_5 is $-C(0)-R_{10}$, wherein R_{10} is Ar_3 and the Ar_3 cyclic group is phenyl, substituted by

O / \ CH₂

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Preferred compounds of this more preferred embodiment include, but are not limited to:

- 220 -

Other compounds of embodiment (K) include,

5 but are not limited to:

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- 227 -

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The ICE inhibitors of another embodiment (L) of this invention are those of formula :

- 231 -

wherein:

m is 1 or 2;

5 R_1 is selected from the group consisting of the following formulae:

C is a ring chosen from the set consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl, the ring optionally being singly or multiply substituted by -Q1;

 R_3 is selected from the group consisting of:

-CN,

-C(O)-H,

-C(O)-CH2-T1-R11,

-C(O)-CH2-F,

 $-C=N-O-R_{Q}$, and

-CO-Arg;

each R_{S} is independently selected from the group consisting of:

-C(O)-R10,

25 -C(0)0-Ra,

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- 232 -

```
-C(0)-N(R10)(R10)
                   -S(O)2-Rg,
                   -S(0)2-NH-R10,
                   -C(0)-CH2-O-R9,
 5
                   -C(0)C(0)-R10.
                   -R<sub>9</sub>.
                   -н.
                   -C(0)C(0)-OR_{10}, and
                   -C(0)C(0)-N(R9)(R10);
1.0
             each T_1 is independently selected from the group
       consisting of -O-, -S-, -S(0)-, and -S(0)<sub>2</sub>-;
            R_6 is selected from the group consisting of -H and
15
       -CH3;
```

```
\ensuremath{R_{\textrm{R}}} is selected from the group consisting of:
                    -C(0)-R<sub>10</sub>,
                    -C(0)0-Rg,
                    -C(0)-NH-R10,
20
                    -S(0)2-Rg,
                    -S (O) 2-NH-R10,
                    -C(O)-CH2-OR10,
                    -C(0)C(0)-R10,
                    -C(0)-CH_2-N(R_{10})(R_{10}),
25
                    -C(O)-CH2C(O)-C-R9,
                    -C(O)-CH2C(O)-R9,
                    -H, and
                    -C(O)-C(O)-OR10;
```

each $R_{\boldsymbol{\mathsf{q}}}$ is independently selected from the group 30 consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar3, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

- 233 -

each R_{10} is independently selected from the group consisting of $^{-}$ H, $^{-}$ Ar $_3$, a C_{3-6} cycloalkyl group, and a $^{-}$ Cl $_{1-6}$ straight or branched alkyl group optionally substituted with Ar $_3$, wherein the $^{-}$ Cl $_{1-6}$ alkyl group is optionally unsaturated;

each \mathbf{R}_{11} is independently selected from the group consisting of:

-Ar₄, - (CH₂)₁₋₃-Ar₄, -H, and -C(O)-Ar₄;

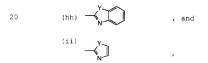
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 R_{15} is selected from the group consisting of -OH, -OAr_3, -N(H)-OH, and -OC_{1-6}, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with Ar_3 , -CONH_2, -OR_5, -OH, -OR_5, or -CO_2H;

Ar₂ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by Q_1 :



wherein each Y is independently selected from the group consisting of O and S;

25 each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains

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6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

each Ar_4 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by - Q_1 ;

each Q1 is independently selected from the group consisting of -NH2, -CO_2H, -Cl, -F, -Br, -I, -NO_2, -CN, =0, -OH, -perfluoro C1_3 alkyl, R5, -OR5, -NHR5, -OR9, -N (R9) (R10^, -R9, -C(0)-R10, and 0 /\ CH2; \ / /

provided that when -Ar $_3$ is substituted with a $\rm Q_1$ group which comprises one or more additional -Ar $_3$

- 235 -

groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Preferably,

m is 1;

5 C is a ring chosen from the set consisting of benzo, pyrido, and thieno, the ring optionally being singly or multiply substituted by halogen, -NH₂, -NH-R₅, or -NH-R₉, -OR₁₀, or -R₉, wherein R₉ is a straight or branched -C₁₋₄ alkyl group, and R₁₀ is -H or a straight or branched -C₁₋₄ alkyl group;

 T_1 is 0 or S;

R6 is H;

 $$\rm R_{11}$ is selected from the group consisting of -Ar4, 15 - (CH2)_1-3-Ar4, and -C(O)-Ar4;

Aro is (hh);

Y is 0;

each Ar₃ cyclic group is independently selected
from the set consisting of phenyl, naphthyl, thienyl,
quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl,
thienothienyl, thiadiazolyl, benzotriazolyl,
benzo(b]thiophenyl, benzofuranyl, and indolyl, and said
cyclic group optionally being singly or multiply
substituted by -0₁;
substituted by -0₁;

each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl,

- 236 -

naphthyl, pyridinyl, oxazolyl, pyrimidinyl, or indolyl, and said cyclic group optionally being singly or multiply substituted by -0_1 ;

each O_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -NH₉, and

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

Preferred compounds of this preferred embodiment include, but are not limited to:

- 237 -

- 238 -

5 711 , and

More preferably, R_3 is -C(0)-Ar $_2$ and the other substituents are as described above. $Alternatively,\ R_3 \ is$

5 -C(0)CH2-T1-R11;

Alternatively, R3 is -C(0)-H.

Preferably, in any of the above compounds of embodiment (L), R_{θ} is selected from the group consisting of:

10 -C(O)-R₁₀,

-C(0)0-Rq,

 $-C(0)-CH_2-OR_{10}$, and

-C(0)-CH₂C(0)-R₉.

More preferably, R_8 is $-\mathrm{C}\left(\mathrm{O}\right)-\mathrm{CH}_2-\mathrm{OR}_{10}$ and

15 R_{10} is -H or -CH₃.

 $\label{eq:alternatively, ICE inhibitors of embodiment} \mbox{(L) of this invention are those of formula :}$

$$(V)$$
 $(I)_{m}$ R_{1}

wherein:

20 m is 1;

R₁ is:

- 240 -

(e10-B)
$$R_{21}$$
 N

 $\ensuremath{\mathrm{R}}_3$ is selected from the group consisting of: -CN,

-C(O)-H,

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 $-C(0)-CH_2-T_1-R_{11}$,

 $-C(0)-CH_2-F$,

-C=N-O-R $_9$, and

-CO-Ar₂;

10 each R_5 is independently selected from the group consisting of:

-C(O)-R₁₀,

-C(0)0-Rg,

-C(0)-N(R₁₀)(R₁₀)

15 -S(0)₂-R₉,

-S(C)2-NH-R₁₀,

-C(O)-CH2-O-R9,

-C(0)C(0)-R₁₀

-R₉,

-H,

-C(O)C(O)-OR $_{10}$, and

-C(0)C(0)-N(Rq)(R10);

 Y_2 is H_2 or O;

25 each T_1 is independently selected from the group consisting of -O- or -S-;

each R_9 is independently selected from the group

- 241 -

consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3 , wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a C_{3-6} cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 $\mbox{ each R_{11} is independently selected from the group } \\ \mbox{10} \qquad \mbox{ consisting of: } \\$

 R_{15} is selected from the group consisting of -OH, -OAr₃, -N(H)-OH, and -OC₁₋₆, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

20 Ar₂ is:

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wherein Y is O:

each Ar₃ is a cyclic group independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl,

- 242 -

isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each Ar_4 is a cyclic group independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₉, and

O / \ CH₂,

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$;

provided that when:

25 m is 1; R_{15} is -OH; R_{21} is -H; and

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 Y_2 is O and R_3 is -C(O)-H, then R_5 cannot be: -C(O)- R_{10} , wherein R_{10} is -Ar $_3$ and the Ar $_3$ cyclic

- 243 -

group is phenyl, unsubstituted by $-Q_1$, 4-(carboxymethoxy)phenyl, 2-fluorophenyl, 2-pyridyl, N-(4-methylpiperazino)methylphenyl, or

-C(0)-OR9, wherein R9 is -CH2-Ar3, and the Ar3 $\,$ cyclic group is phenyl, unsubstituted by -Q1; and when

 $\rm Y_2$ is 0, $\rm R_3$ is -C(0)-CH₂-T₁-R₁₁, T₁ is 0, and R₁₁ is Ar₄, wherein the Ar₄ cyclic group is 5-(1-(4-chlorophenyl)-3-trifluoromethyl)pyrazolyl), then R₅ cannot be:

10 -H:

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 $-C(0)-R_{10}$, wherein R_{10} is $-Ar_3$ and the Ar_3 cyclic group is 4-(dimethylaminomethyl)phenyl, phenyl, 4-(carboxymethylthio)phenyl, 4-(carboxymethylthio)phenyl, 4-(carboxyethyl)phenyl, 4-(carboxyethyl)phenyl, 2-pyridyl, N-(4-

methylpiperazino)methylphenyl, or

-C(0)-OR9, wherein $\rm R_9$ is isobutyl or -CH_2-Ar_3 and the Ar_3 cyclic group is phenyl;

and when R_{11} is Ar_4 , wherein the Ar_4 cyclic group 20 is $5-(1-phenyl-3-trifluoromethyl)pyrazolyl or <math>5-(1-(4-chloro-2-pyridinyl)-3-trifluoromethyl)pyrazolyl, then <math>R_5$ cannot be:

-C(0)-OR9, wherein R9 is -CH2-Ar3, and the Ar3 cyclic group is phenyl;

and when R_{11} is Ar_4 , wherein the Ar_4 cyclic group is 5-(1-(2-pyridyl)-3-trifluoromethyl)pyrazolyl), then R_5 cannot be:

-C(0)-R $_{10}$, wherein R_{10} is -Ar $_3$ and the Ar $_3$ cyclic group is 4-(dimethylaminomethyl)phenyl, or

-C(0)-OR9, wherein R_9 is -CH2-Ar3, and the Ar3 cyclic group is phenyl, unsubstituted by -O1; and when

- 244 -

 $\rm Y_2$ is 0, $\rm R_3$ is -C(0)-CH₂-T₁-R₁₁, T₁ is 0, and R₁₁ is -C(0)-Ar₄, wherein the Ar₄ cyclic group is 2,5-dichlorophenyl, then R₅ cannot be:

-C(0)-R₁₀, wherein R₁₀ is -Ar₃ and the Ar₃ cyclic group is 4-(dimethylaminomethyl)phenyl, 4-(N-morpholinomethyl)phenyl, 4-(N-methylpiperazino)methyl)phenyl, 4-(N-(2-methyl)imidazolylmethyl)phenyl, 5-benzimidazolyl, 5-benztriazolyl, N-carboethoxy-5-benztriazolyl, N-carboethoxy-5-benztriazolyl, N-carboethoxy-5-benztriazolyl, N-carboethoxy-5-benztriazolyl, N-

-C(0)-OR9, wherein $\rm R_9$ is -CH2-Ar3, and the Ar3 cyclic group is phenyl, unsubstituted by -Q1,; and when

 $\rm Y_2$ is $\rm H_2,~R_3$ is -C(0)-CH₂-T₁-R₁₁, T₁ is 0, and R₁₁ is -C(0)-Ar₄, wherein the Ar₄ cyclic group is 2,5-dichlorophenyl, then R₅ cannot be:

-C(0)-OR9, wherein $\rm R_9$ is -CH2-Ar3 and the $\rm Ar_3$ cyclic group is phenyl.

Preferably, in any of the above compounds of embodiment (L), R $_3$ is -C(0)-H and R $_5$ is -C(0)-R $_{10}$ or -C(0)-C(0)-R $_{10}$ and the other substituents are as defined above.

More preferably $\ensuremath{R_{10}}$ is $-\ensuremath{Ar_3}$ and the other substituents are as defined above.

More preferably in these more preferred

 $\rm R_5$ is -C(O)-R_{10} and R_{10} is Ar_3, wherein the Ar_3 cyclic group is phenyl optionally being singly or multiply substituted by:

 $-R_9$, wherein R_9 is a C_{1-4} straight or branched alkyl group;

-F,

compounds:

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-C1,

 $-N(H)-R_5$, wherein $-R_5$ is -E or $-C(0)-R_{10}$,

- 245 -

wherein $\rm R_{10}$ is a $^{\rm -}C_{1-6}$ straight or branched alkyl group optionally substituted with Ar_3, wherein Ar_3 is phenyl,

 $^{-N}(\rm R_{9})\;(R_{10})\,,$ wherein $\rm R_{9}$ and $\rm R_{10}$ are independently a $^{-C}_{1-4}$ straight or branched alkyl group, or

-O-R5, wherein R5 is H or a -C1-4 straight or branched alkyl group.

Preferred compounds of this preferred embodiment include, but are not limited to:

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Most preferably, Ar $_3$ is phenyl being singly or multiply substituted at the 3- or 5-position by -Cl or at the 4-position by -NH-R $_5$, -N(R $_9$)(R $_{10}$), or -O-R $_5$.

5 Preferred compounds of this most preferred embodiment include, but are not limited to:

- 248 -

- 249 -

Other preferred compounds of this most preferred embodiment include, but are not limited to:

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Alternatively, Ar_3 is phenyl being singly or multiply substituted at the 3- or 5-position by $-R_9$, wherein R_9 is a C_{1-4} straight or branched alkyl group; and at the 4-position by $-O-R_5$.

Preferred compounds of this most preferred embodiment include, but are not limited to:

- 250 -

Another preferred compound of

5 this most preferred embodiment includes, but is not limited to:

- 252 -

Alternatively, in this more preferred

embodiment:

 $\rm R_5$ is -C(0)-R₁₀, wherein R₁₀ is Ar₃ and the Ar₃ cyclic group is selected from the group consisting of is indolyl, benzimidazolyl, thienyl, quinolyl, isoquinolyl and benzo(b)thiophenyl, and said cyclic group optionally being singly or multiply substituted by -Q₁.

Preferred compounds of this more preferred embodiment include, but are not limited to:

Most preferably, the Ar_3 cyclic group is isoquinoly!, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.

A preferred compound of this most preferred embodiment includes, but is not limited

- 253 -

to:

696-1 ; and

Another preferred compound of this most preferred embodiment includes, but is not limited to:

- 254 -

Alternatively, in this more preferred embodiment R_5 is $-C(0)-R_{10}$, wherein R_{10} is $-Ar_3$ and the Ar $_3$ cyclic group is phenyl, substituted by

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A preferred compound of this more preferred embodiment includes, but is not limited to:

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A preferred compound of this more preferred embodiment includes, but is not limited to:

Other compounds of embodiment $(\ensuremath{\mathrm{L}})$ include, but are not limited to:

- 258 -

- 259 -

- 260 -

- 262 -

- 263 -

5 708 , N N N H

- 265 -

- 266 -

- 267 -

- 268 -

- 269 -

Other compounds of embodiment $(\ensuremath{\mathrm{K}})$ include, but are not limited to:

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- 271 -

Other compounds of embodiment $(\ensuremath{\mathrm{L}})$ include, but are not limited to:

- 274 -

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- 276 -

- 277 -

- 278 -

744

746

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- 281 -

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- 282 -

- 284 -

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- 285 -

The most preferred compounds of embodiments $(\mbox{\it K})$ and $(\mbox{\it L})$ are those wherein the $\mbox{\it Ar}_3$ cyclic group is isoquinolyl.

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1.5

2.0

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Compounds of this invention are described in co-pending United States Application Serial Nos. 08/575,641 and 08/598,332 the disclosures of which are herein incorporated by reference.

The compounds of this invention have a molecular weight of less than or equal to about 700 Daltons, and more preferably between about 400 and 600 Daltons. These preferred compounds may be readily absorbed by the bloodstream of patients upon oral administration. This oral availability makes such compounds excellent agents for orally-administered treatment and prevention regimens against IL-1-, apoptosis-, IGIF- or IFN-y mediated diseases.

It should be understood that the compounds of this invention may exist in various equilibrium forms, depending on conditions including choice of solvent, pH, and others known to the practitioner skilled in the art. All such forms of these compounds are expressly included in the present invention. In particular, many

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of the compounds of this invention, especially those which contain aldehyde or ketone groups in R_3 and carboxylic acid groups in T, may take hemi-ketal (or hemi-acetal) or hydrated forms. For example, compounds of embodiment (A) may take the forms depicted below: EO1

Depending on the choice of solvent and other conditions known to the practitioner skilled in the art, compounds of this invention may also take acyloxy ketal, acyloxy acetal, ketal or acetal form:

In addition, it should be understook that the equilibrium forms of the compounds of this invention may include tautomeric forms. All such forms of these compounds are expressly included in the present invention.

It should be understood that the compounds of this invention may be modified by appropriate

- 287 -

functionalities to enhance selective biological properties. Such modifications are known in the art and include those which increase biological penetration into a given biological system (e.g., blood, lymphatic 5 system, central nervous system), increase oral availability, increase solubility to allow administration by injection, alter metabolism and alter rate of excretion. In addition, the compounds may be altered to pro-drug form such that the desired compound 1.0 is created in the body of the patient as the result of the action of metabolic or other biochemical processes on the pro-drug. Such pro-drug forms typically demonstrate little or no activity in in vitro assays. Some examples of pro-drug forms include ketal, acetal, 15 oxime, imine, and hydrazone forms of compounds which contain ketone or aldehyde groups, especially where they occur in the R3 group of the compounds of this invention. Other examples of pro-drug forms include the hemi-ketal, hemi-acetal, acyloxy ketal, acyloxy 20 acetal, ketal, and acetal forms that are described in EO1 and EO2.

ICE and TX Cleave and Thereby Activate Pro-IGIF

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The ICE protease was identified previously by virtue of its ability to process inactive pro-IL-18 to mature active IL-18, a pro-inflammatory molecule, in virro and in vivo. Here we show that ICE and its close homologue TX (Caspase-4, C. Faucheu et al., EMBO, 14, p. 1914 (1995)) can proteolytically cleave inactive pro-IGIF. This processing step is required to convert pro-IGIF to its active mature form, IGIF. Cleavage of pro-IGIF by ICE, and presumably by TX, also facilitates the export of IGIF our of cells.

- 288 -

We first used transient co-expression of plasmids transfected into Cos cells to determine whether any known members of the ICE/CED-3 protease family can process pro-IGIF to IGIF in cultured cells (Example 23) (Fig. 1A).

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Fig. 1A demonstrates that ICE cleaves pro-IGIF in Cos cells co-transfected with plasmids that express pro-IGIF in the presence of active ICE. Cos cells were transfected with an expression plasmid for pro-IGIF alone (lane 2) or in combination with the indicated expression plasmids encoding wild type or inactive mutants of ICE/CED-3 family of proteases (lanes 3-12). Cell lysates were prepared and analyzed for the presence of IGIF protein by immunoblotting with an anti-IGIF antiserum. Lane 1 contained lysates from mock transfected cells.

Co-expression of pro-IGIF with ICE or TX resulted in the cleavage of pro-IGIF into a polypeptide similar in size to the naturally-occurring 18-kDa mature IGIF. This processing event is blocked by single point mutations that alter the catalytic cysteine residues and thus inactivate ICE and TX (Y. Gu et al., EMBO, 14, p. 1923 (1995)).

Co-expression with CPP32 (Caspase-3), a

25 protease involved in programmed cell death (T.
Fernandes-Alnemri et al., J. Biol. Chem., 269, p. 30761
(1994); D. W. Nicholson et al., Nature, 376, p. 37
(1995)), resulted in the cleavage of pro-IGIF into a
smaller polypeptide, while co-expression with CMH-1

30 (Caspase-7), a close homolog of CPP32 (J. A. Lippke et
al., J. Biol. Chem., 271, p. 1825 (1996)), failed to
cleave pro-IGIF to any significant extent. Thus, ICE
and TN appear to be capable of cleaving pro-IGIF into a

- 289 -

polypeptide similar in size to the naturally-occurring 18-kDa IGIF.

We next examined the ability of these cysteine proteases to cleave pro-IGIF <u>in vitro</u> using a purified, recombinant (His)₆-tagged pro-IGIF as a substrate (Example 23).

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Fig. 1B demonstrates that pro-IGIF is cleaved in vitro by ICE. Purified recombinant (His).-tagged pro-IGIF (2 μg) was incubated with the indicated cysteine protease in the presence or absence of ICE or CPP32 inhibitors as described in Example 23. The cleavage products were analyzed by SDS-PAGE and Coomassie Blue staining.

ICE cleaved the 24 kDa pro-IGIF into two
polypeptides of approximately 18-kDa and 6-kDa.

N-terminal amino acid sequencing of the ICE cleavage
products indicated that the 18-kDa polypeptide contains
the same N-terminal amino acid residues
(Asn-Phe-Gly-Arg-Leu) as the naturally occurring IGIF.
This shows that ICE cleaves pro-IGIF at the authentic
processing site (Asp35-Asn36) (H. Okamura et al.,
Infection and Immunity, 63, p. 3966 (1995); H. Okamura
et al., Nature, 378, p. 88 (1995)). N-terminal amino
acid sequencing of the CPP32 cleavage products

indicated that CPP32 cleaved pro-IGIF at Asp69-Ile70. The cleavage by ICE of pro-IGIF is highly specific with a catalytic efficiency (k_{cat}/K_M) of i.4 x 10^7 m⁻¹ s⁻¹ $(K_M=0.6\pm0.1~\mu\text{M};~k_{cat}=8.6\pm0.3~\text{s}^{-1})$ and is inhibited by specific ICE inhibitors

30 (Ac-Tyr-Val-Ala-Asp-aldehyde) and Cbz-Val-Ala-Asp[:2,6-dichlorobenzoyl)oxy]methylketone, (N.A.
Thornberry et al., Nature, 356, p. 768 (1992); R. E.
Dolle et al., J. Med. Chem., 37, p. 563 (1994)}.

- 290 -

Fig. 1C demonstrates that ICE cleavage in vitro activates pro-IGIF. Uncleaved pro-IGIF, ICE- or CPP32-cleaved products of pro-IGIF, or recombinant mature IGIF (rIGIF) were each added to A.E7 cell cultures to a final concentration of 12 ng/ml or 120 ng/ml (see, Example 23). Eighteen hours later, IFN-y in the cultural medium was quantified by ELISA. While the uncleaved pro-IGIF had no detectable IFN-y inducing activity, ICE-cleaved pro-IGIF was active in inducing IFN-y production in Thl cells.

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3.0

Like ICE, the ICE homolog TX also cleaved pro-IGIF into similarly sized polypeptides. However, its catalytic efficiency was about two orders of magnitude lower than that shown for ICE.

Consistent with the observations from the Cos cell experiments above, CPP32 cleaved pro-IGIF at a different site (Asp69-Ile70) and the resulting polypeptides had little IFN-y inducing activity (Fig. 1C). CMH-1 and granzyme B each failed to cleave pro-IGIF to any significant extent.

Together, these results demonstrate that, both in Cos cells and in vitro, ICE and TX are capable of processing the inactive pro-IGIF precursor at the authentic maturation site to generate a biologically active IGIF molecule.

Processing of Pro-IGIF by ICE Pacilitates Its Export

IGIF is produced by activated Kupffer cells
and macrophages in vivo and is exported out of the
cells upon stimulation by endotoxin (H. Okamura et al.,
Infection and Immunity, 63, p. 3966 (1995); H. Okamura
et al., Nature, 378, p. 88 (1995). We used the Cos
cell co-expression system (Example 23) to examine

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whether the intracellular cleavage of pro-IGIF by ICE would facilitate the export of mature IGIF from the cell. Such is the case for pro-IL-1 β when it is cleaved by ICE into active IL-1 β (N.A. Thornberry et al., Nature, 356, p. 768 (1992)).

In Fig. 2A, Cos cells transfected with an expression plasmid for pro-IGIF alone (lanes 2 and 6) or in combination with an expression plasmid encoding wild type (lanes 3 and 7) or inactive mutant ICE (lanes 4 and 8) were metabolically labeled with ³⁵S-methionine (see, Example 24). Cell lysates (left) and conditioned medium (right) were immunoprecipitated with an anti-IGIF antiserum. The immunoprecipitated proteins were analyzed by SDS-PAGE and fluorography (Fig. 2A).

An 18-kDa polypeptide corresponding in size to mature IGIF was detected in the conditioned medium of Cos cells co-expressing pro-IGIF and ICE, while Cos cells co-expressing pro-IGIF and an inactive ICE mutant (ICE-C285S), or pro-IGIF alone (-) exported only very low levels of pro-IGIF and no detectable mature IGIF. We estimate that about 10% of the mature IGIF was exported from co-transfected cells, while greater than 99% of pro-IGIF was retained within the cells.

We also measured the presence of IFN-y
inducing activity in cell lysates and in the
conditioned medium of the above transfected cells (see,
Example 24). IFN-y inducing activity was detected in
both cell lysates and the conditioned medium of Cos
cells co-expressing pro-IGIF and ICE, but not in cells
expressing either pro-IGIF or ICE alone (Fig. 2B.

These results indicate that ICE cleavage of pro-IGIF facilitates the export of mature, active IGIF from cells.

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Pro-IGIF is a Physiological Substrate of ICE In Vivo

To study the role of ICE in the proteolytic activation and export of IGIF under physiological conditions, we examined the processing of pro-IGIF and export of mature IGIF from lipopolysaccharide (LPS)-activated Kupffer cells harvested from

(LPS)-activated Kupffer cells harvested from <u>Propiobacterium acnes</u>-elicited wild type and ICE deficient (ICE-/-) mice (Example 25).

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As shown in Fig. 3A, Kupffer cells from

ICE-/- mice are defective in the export of IGIF.

Kupffer cell lysates of wild type and ICE-/- mice
contained similar amounts of IGIF as determined by
ELISA. IGIF, however, could be detected only in the
conditioned medium of wild type but not of the ICE-/
cells. Thus, ICE-deficient (ICE-/-) mice synthesize
pro-IGIF, but fail to export it as extracellular pro-or
mature IGIF.

To determine whether ICE-deficient (ICE-/-)
mice process intracellular pro-IGIF but fail to export
IGIF, Kupffer cells from wild type and ICE-/- mice were
metabolically labeled with "S-methionine and IGIF
immunoprecipitation experiments were performed on cell
lysates and conditioned media as described in Example
25. These experiments demonstrated that unprocessed
pro-IGIF was present in both wild type and ICE-/-

Kupffer cells. However, the 18-kDa mature IGIF was present only in the conditioned medium of wild type and not ICE-/- Kupffer cells (Fig. 3B). This shows that active ICE is required in cells for the export of processed IGIF out of the cell.

In addition, conditioned medium from wild type but not from ICE-/- Kupffer cells contained IFN- γ inducing activity that was not attributed to the action

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of IL-12 because it was insensitive to a neutralizing anti-IL-12 antibody. The absence of IGIF in the conditioned medium of ICE-/- Kupffer cells is consistent with the finding in Cos cells that the processing of pro-IGIF by ICE is required for the export of active IGIF.

Figs. 3C and 3D show that, in vivo, ICE-/mice have reduced serum levels of IGIF and IFN-y,
respectively. Wild type (ICE+/+) and ICE-/- mice (n=3)
primed with heat-inactivated P. acnes were challenged
with LPS (Example 26), and the levels of IGIF (Fig. 3C)
and IFN-y (Fig. 3D) in the sera of challenged mice were
measured by ELISA three hours after LPS challenge
(Example 25).

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15 The sera of ICE-/- mice stimulated by P. acnes and LPS contained reduced levels of IGIF (Fig. 3C) and no detectable IFN- γ inducing activity in the presence of an anti-IL-12 antibody. The reduced serum levels of IGIF likely accounts for the 20 significantly lower levels of IFN-y in the sera of ICE-/- mice (Fig. 3D), because we have observed no significant difference in the production of IL-12 in ICE-/- mice under these conditions. Consistent with this interpretation is the finding that non-adherent 25 splenocytes from wild type and ICE-/- mice produced similar amounts of IFN-y when stimulated with recombinant active IGIF in vitro. Thus the impaired production of IFN-y is not due to any apparent defect in the T cells of the ICE-/- mice.

Taken together, these results establish a critical role for ICE in processing the IGIF precursor and in the export of active IGIF both in vitro and in

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vivo.

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To examine in more detail the relationship between serum levels of IFN- γ and ICE activity in vivo, a time course after challenge of wild type and ICE-deficient mice with LPS was performed (Example 26) (Fig. 4).

Fig. 4 shows a time course increase of serum IFN- γ in wild type mice, with sustained levels of \$\grel17\$ ng/ml occurring from 9-18 hrs after LPS challenge. As predicted by the experiments discussed above, serum IFN- γ levels were significantly lower in ICE-/- mice, with a maximum of 2 ng/ml achieved over the same time period, which is approximately 15% of the level observed in wild type mice (Fig. 4).

Animals were also observed for clinical signs of sepsis and body temperature was measured at 4-hour intervals in wild type and ICE-/- mice challenged with 30 mg/kg or 100 mg/kg LPS (ICE-/-only). Results in Fig. 4 show that wild type mice experienced a significant decrease in body temperature (from 36°C to 26°C) within 12 hours of LPS challenge. Signs of clinical sepsis were evident and all animals expired within 24-28 hours.

In contrast, ICE-/- mice challenged with

30 mg/kg LPS experienced only a 3°-4°C decrease in body
temperature with minimal signs of distress and with no
observed lethality. ICE-/- mice challenged with
100 mg/kg LPS experienced clinical symptoms, a decrease
in body temperature, and mortality similar to wild type
30 mice at the 30 mg/kg LPS dose.

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The ICE Inhibitor Ac-YVAD-CHO is an Equipotent Inhibitor of IL-18 and IFN-y Production

Since the processing and secretion of biologically active IGIF is mediated by ICE, we compared the activity of a reversible ICE inhibitor (Ac-YVAD-CHO) on IL-1 β and IFN- γ production in a peripheral blood mononuclear cell (PBMC) assay (Examples 27).

Results in Fig. 5 show a similar potency for the ability of the Ac-YVAD-CHO ICE inhibitor to decrease IL-1 β and IFN- γ production in human FBMCs, with an IC50 of 2.5 μ M for each. Similar results were obtained in studies with wild type mouse splenocytes. These findings provide additional evidence that pro-IGIF is a physiological substrate for ICE and suggest that ICE inhibitors will be useful tools for

In summary, we have found that ICE controls IGIF and IFN- γ levels in vivo and in vitro and that ICE inhibitors can decrease levels of IGIF and IFN- γ in human cells. These results have been described in copending United States Application Serial No. 08/712,878, the disclosure of which is herein incorporated by reference.

controlling physiological levels of IGIF and IFN-y.

Compositions and Methods

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The pharmaceutical compositions and methods of this invention will be useful for controlling IL-1, IGIF and IFN-y levels in vivo. The methods and compositions of this invention will thus be useful for treating or reducing the advancement, severity of effects of IL-1, IGIF- and IFN-y-mediated conditions.

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The compounds of this invention are effective ligands for ICE. Accordingly, these compounds are capable of targeting and inhibiting events in IL-1-. apoptosis-, IGIF-, and IFN-y-mediated diseases, and, 5 thus, the ultimate activity of that protein in inflammatory diseases, autoimmune diseases, destructive bone, proliferative disorders, infectious diseases, and degenerative diseases. For example, the compounds of this invention inhibit the conversion of precursor IL- 1β to mature IL- 1β by inhibiting ICE. Because ICE is 1.0 essential for the production of mature IL-1, inhibition of that enzyme effectively blocks initiation of IL-1mediated physiological effects and symptoms, such as inflammation, by inhibiting the production of mature 1.5 IL-1. Thus, by inhibiting IL-1 β precursor activity, the compounds of this invention effectively function as IL-1 inhibitors.

Similarly, compounds of this invention inhibit the conversion of precursor IGIF to mature 20 IGIF. Thus, by inhibiting IGIF production, the compounds of this invention effectively function as inhibitors of IFN-y production.

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Accordingly, one embodiment of this invention provides a method for decreasing IGIF production in a subject comprising the step of administering to the subject a pharmaceutical composition comprising a therapeutically effective amount of an ICE inhibitor and a pharmaceutically acceptable carrier.

Another embodiment of this invention provides

a method for decreasing IFN-y production in a subject
comprising the step of administering to the subject a
pharmaceutical composition comprising a therapeutically
effective amount of an ICE inhibitor and a

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pharmaceutically acceptable carrier.

administering a TX inhibitor.

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In another embodiment, the methods of this invention comprise the step of administering to a subject a pharmaceutical composition comprising an inhibitor of an ICE-related protease that is capable of cleaving pro-IGIF to active IGIF, and a pharmaceutically acceptable carrier. One such ICE-related protease is TX, as described above. This invention thus provides methods and pharmaceutical compositions for controlling IGIF and IFN-Y levels by

Other ICE-related proteases capable of processing pro-IGIF into an active IGIF form may also be found. Thus it is envisioned that inhibitors of those enzymes may be identified by those of skill in the art and will also fall within the scope of this invention.

The compounds of this invention may be employed in a conventional manner for the treatment of diseases which are mediated by IL-1, apoptosis, IGIF or IFN-y. Such methods of treatment, their dosage levels and requirements may be selected by those of ordinary skill in the art from available methods and techniques. For example, a compound of this invention may be combined with a pharmaceutically acceptable adjuvant for administration to a patient suffering from an IL-1-, apoptosis-, IGIF- or IFN-y-mediated disease in a pharmaceutically acceptable manner and in an amount effective to lessen the severity of that disease.

Alternatively, the compounds of this invention may be used in compositions and methods for treating or protecting individuals against IL-1-, apoptosis-, IGIF- or IFN-y-mediated diseases over

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extended periods of time. The compounds may be employed in such compositions either alone or together with other compounds of this invention in a manner consistent with the conventional utilization of ICE inhibitors in pharmaceutical compositions. For example, a compound of this invention may be combined with pharmaceutically acceptable adjuvants conventionally employed in vaccines and administered in prophylactically effective amounts to protect individuals over an extended period of time against IL-1-, apoptosis-, IGIF- or IFN-y- mediated diseases.

The compounds of this invention may also be co-administered with other ICE inhibitors to increase the effect of therapy or prophylaxis against various

In addition, the compounds of this invention may be used in combination either conventional anti-inflammatory agents or with matrix metalloprotease inhibitors, lipoxygenase inhibitors and antagonists of cytokines other than ID-18.

IL-1-, apoptosis, IGIF- or IFN-y-mediated diseases.

The compounds of this invention can also be administered in combination with immunomodulators (e.g., bropirimine, anti-human alpha interferon antibody, IL-2, GM-CSF, methionine enkephalin, interferon alpha, diethyldithiocarbamate, tumor necrosis factor, naltrexone and rEPO) or with prostaglandins, to prevent or combat IL-1-mediated disease symptoms such as inflammation.

When the compounds of this invention are administered in combination therapies with other agents, they may be administered sequentially or concurrently to the patient. Alternatively, pharmaceutical or prophylactic compositions according

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to this invention comprise a combination of an ICE inhibitor of this invention and another therapeutic or prophylactic agent.

Pharmaceutical compositions of this invention 5 comprise any of the compounds of the present invention, and pharmaceutically acceptable salts thereof, with any pharmaceutically acceptable carrier, adjuvant or vehicle. Pharmaceutically acceptable carriers. adjuvants and vehicles that may be used in the pharmaceutical compositions of this invention include, 10 but are not limited to, ion exchangers, alumina, aluminum stearate, lecithin, self-emulsifying drug delivery systems (SEDDS) such as dα-tocopherol polyethyleneglycol 1000 succinate, or other similar 15 polymeric delivery matrices, serum proteins, such as human serum albumin, buffer substances such as phosphates, glycine, sorbic acid, potassium sorbate, partial glyceride mixtures of saturated vegetable fatty acids, water, salts or electrolytes, such as protamine 20 sulfate, disodium hydrogen phosphate, potassium hydrogen phosphate, sodium chloride, zinc salts. colloidal silica, magnesium trisilicate, polyvinyl pyrrolidone, cellulose-based substances, polyethylene glycol, sodium carboxymethylcellulose, polyacrylates, 25 waxes, polyethylene-polyoxypropylene-block polymers, polyethylene glycol and wool fat. Cyclodextrins such as α -, β - and γ -cyclodextrin, or chemically modified derivatives such as hydroxyalkylcyclodextrins, including 2-and 3-hydroxypropyl-β-cyclodextrines, or 30 other solubiliezed derivatives may also be advantageeously used to enhanve delivery of compounds of this invention.

The pharmaceutical compositions of this invention may be administered orally, parenterally, by

- 300 - inhalation spray, topically, rectally, nasally,

buccally, vaginally or via an implanted reservoir. prefer oral administration. The pharmaceutical compositions of this invention may contain any 5 conventional non-toxic pharmaceutically-acceptable carriers, adjuvants or vehicles. In some cases, the pH of the formulation may be adjusted with pharmaceutically acceptable acids, bases or buffers to enhance the stability of the formulated compounds or 10 its delivery form. The term parenteral as used herein includes subcutaneous, intracutaneous, intravenous, intramuscular, intra-articular, intrasynovial, intrasternal, intrathecal, intralesional and intracranial injection or infusion techniques. 15 The pharmaceutical compositions may be in the form of a sterile injectable preparation, for example, as a sterile injectable aqueous or oleaginous suspension. This suspension may be formulated according to techniques known in the art using suitable 20 dispersing or wetting agents (such as, for example, Tween 80) and suspending agents. The sterile injectable preparation may also be a sterile injectable solution or suspension in a non-toxic parenterallyacceptable diluent or solvent, for example, as a 2.5 solution in 1,3-butanediol. Among the acceptable vehicles and solvents that may be employed are mannitol, water, Ringer's solution and isotonic sodium

medium. For this purpose, any bland fixed oil may be employed including synthetic mono- or diglycerides. Fatty acids, such as oleic acid and its glyceride derivatives are useful in the preparation of injectables, as are natural pharmaceutically-acceptable

chloride solution. In addition, sterile, fixed oils are conventionally employed as a solvent or suspending

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oils, such as olive oil or castor oil, especially in their polyoxyethylated versions. These oil solutions or suspensions may also contain a long-chain alcohol diluent or dispersant such as <u>Ph</u>, <u>Helv</u> or a similar alcohol.

The pharmaceutical compositions of this invention may be orally administered in any orally acceptable dosage form including, but not limited to, capsules, tablets, and aqueous suspensions and solutions. In the case of tablets for oral use, carriers which are commonly used include lacticse and corn starch. Lubricating agents, such as magnesium stearate, are also typically added. For oral administration in a capsule form, useful diluents include lactose and dried corn starch. When aqueous suspensions are administered orally, the active ingredient is combined with emulsifying and suspending agents. If desired, certain sweetening and/or flavoring and/or coloring agents may be added.

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The pharmaceutical compositions of this invention may also be administered in the form of suppositories for rectal administration. These compositions can be prepared by mixing a compound of this invention with a suitable non-irritating excipient which is solid at room temperature but liquid at the rectal temperature and therefore will melt in the rectum to release the active components. Such materials include, but are not limited to, cocoa butter, beeswax and polyethylene glycols.

Topical administration of the pharmaceutical compositions of this invention is especially useful when the desired treatment involves areas or organs readily accessible by topical application. For application topically to the skin, the pharmaceutical

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composition should be formulated with a suitable ointment containing the active components suspended or dissolved in a carrier. Carriers for topical administration of the compounds of this invention . include, but are not limited to, mineral oil, liquid petroleum, white petroleum, propylene glycol, polyoxyethylene polyoxypropylene compound, emulsifying wax and water. Alternatively, the pharmaceutical composition can be formulated with a suitable lotion or cream containing the active compound suspended or dissolved 10 in a carrier. Suitable carriers include, but are not limited to, mineral oil, sorbitan monostearate, polysorbate 60, cetyl esters wax, cetearyl alcohol, 2-octyldodecanol, benzyl alcohol and water. The 15 pharmaceutical compositions of this invention may also be topically applied to the lower intestinal tract by rectal suppository formulation or in a sustable enema formulation. Topically-administered transdermal patches are also included in this invention. 20

The pharmaceutical compositions of this invention may be administered by nasal aerosol or inhalation. Such compositions are prepared according to techniques well-known in the art of pharmaceutical formulation and may be prepared as solutions in saline, employing benzyl alcohol or other suitable preservatives, absorption promoters to enhance bloavailability, fluorocarbons, and/or other solubilizing or dispersing agents known in the art.

Dosage levels of between about 0.01 and about

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30 100 mg/kg body weight per day, preferably between about 1 and 50 mg/kg body weight per day of the active ingredient compound are useful in the prevention and treatment of IL-1-, apoptosis, IGIF and IFN-y-mediated

diseases, including inflammatory diseases, autoimmune diseases, destructive bone disorders, proliferative disorders, infectious diseases, degenerative diseases. necrotic diseases, osteoarthritis, acute pancreatitis, 5 chronic pancreatitis, asthma, adult respiratory distress syndrome, glomeralonephritis, rheumatoid arthritis, systemic lupus erythematosus, scleroderma, chronic thyroiditis, Graves' disease, autoimmune gastritis, insulin-dependent diabetes mellitus (Type 1.0 I), autoimmune hemolytic anemia, autoimmune neutropenia, thrombocytopenia, chronic active hepatitis, myasthenia gravis, inflammatory bowel disease, Crohn's disease, psoriasis, graft vs. host disease, osteoporosis, multiple myeloma-related bone 15 disorder, acute myelogenous leukemia, chronic mvelogenous leukemia, metastatic melanoma, Kaposi's sarcoma, multiple myeloma sepsis, septic shock, Shigellosis, Alzheimer's disease, Parkinson's disease, cerebral ischemia, myocardial ischemia, spinal muscular atrophy, multiple sclerosis, AIDS-related encephalitis, 20 HIV-related encephalitis, aging, alopecia, and neurological damage due to stroke. Typically, the pharmaceutical compositions of this invention will be administered from about 1 to 5 times per day or 25 alternatively, as a continuous infusion. Such administration can be used as a chronic or acute therapy. The amount of active ingredient that may be combined with the carrier materials to produce a single dosage form will vary depending upon the host treated and the particular mode of administration. A typical 30 preparation will contain from about 5% to about 95; active compound (w/w). Preferably, such preparations contain from about 20% to about 80% active compound. Upon improvement of a patient's condition, a

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maintenance dose of a compound, composition or combination of this invention may be administered, if necessary. Subsequently, the dosage or frequency of administration, or both, may be reduced, as a function of the symptoms, to a level at which the improved condition is retained when the symptoms have been alleviated to the desired level, treatment should cease. Patients may, however, require intermittent treatment on a long-term basis upon any recurrence or disease symptoms.

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As the skilled artisan will appreciate, lower or higher doses than those recited above may be required. Specific dosage and treatment regimens for any particular patient will depend upon a variety of factors, including the activity of the specific compound employed, the age, body weight, general health status, sex, diet, time of administration, rate of excretion, drug combination, the severity and course of the disease, and the patient's disposition to the disease and the judgment of the treating physician.

The IL-1 mediated diseases which may be treated or prevented by the compounds of this invention include, but are not limited to, inflammatory diseases, autoimmune diseases, destructive bone disorders, proliferative disorders, infectious diseases, and degenerative diseases. The apoptosis-mediated diseases which may be treated or prevented by the compounds of

Inflammatory diseases which may be treated or prevented include, but are not limited to osteoarthritis, acute pancreatitis, chronic pancreatitis, asthma, and adult respiratory distress syndrome. Preferably the inflammatory disease is osteoarthritis or acute pancreatitis.

this invention include degenerative diseases.

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Autoimmune diseases which may be treated or prevented include, but are not limited to, glomeralonephritis, rheumatoid arthritis, systemic lupus erythematosus, scleroderma, chronic thyroiditis, Graves' disease, autoimmune gastritis, insulindependent diabetes mellitus (Type I), autoimmune hemolytic anemia, autoimmune neutropenia, thrombocytopenia, chronic active hepatitis, myasthenia gravis, multiple sclerosis, inflammatory bowel disease, Crohn's disease, psoriasis, and graft vs. host disease. Preferably the autoimmune disease is rheumatoid arthritis, inflammatory bowel disease, Crohn's disease,

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or psoriasis.

Destructive bone disorders which may be

treated or prevented include, but are not limited to,
osteoporosis and multiple myeloma-related bone
disorder.

Proliferative diseases which may be treated or prevented include, but are not limited to, acute myelogenous leukemia, chronic myelogenous leukemia, metastatic melanoma, Kaposi's sarcoma, and multiple myeloma.

Infectious diseases which may be treated or prevented include, but are not limited to, sepsis, septic shock, and Shigellosis.

The IL-1-mediated degenerative or necrotic diseases which may be treated or prevented by the compounds of this invention include, but are not limited to, Alzheimer's disease, Parkinson's disease, cerebral ischemia, and myccardial ischemia.

Preferably, the degenerative disease is Alzheimer's disease

The apoptosis-mediated degenerative diseases which may be treated or prevented by the compounds of

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this invention include, but are not limited to, Alzheimer's disease, Parkinson's disease, cerebral ischemia, myocardial ischemia, spinal muscular atrophy, multiple sclerosis, AIDS-related encephalitis, HIV-related encephalitis, aging, alopecia, and neurological damage due to stroke.

The methods of this invention may be used for treating, or reducing the advancement, severity or effects of an IGIF-or IFN- γ -mediated inflammatory, autoimmune, infectious, proliferative, destructive bone, necrotic, and degenerative conditions, including

diseases, disorders or effects, wherein the conditions are characterized by increased levels of IGIF or IFN- γ production.

production.

Examples of such inflammatory conditions include, but are not limited to, osteoarthritis, acute pancreatitis, chronic pancreatitis, asthma, rheumatord arthritis, inflammatory bowel disease, Crohn's disease, ulcerative collitis, cerebral ischemia, myocardial ischemia and adult respiratory distress syndrome.

Preferably, the inflammatory condition is rheumatoid arthritis, ulcerative collitis, Crohn's disease, hepatitis and adult respiratory distress syndrome.

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Examples of such infectious conditions include, but are not limited to, infectious hepatitis, sepsi~ septic shock and Shigellosis.

Examples of such autoimmune conditions incluie, but are not limited to, glomerulonephritis, systemic lupus erythematosus, scleroderma, chronic thyroiditis, Graves' disease, autoimmune gastritis, insulin-dependent diabetes mellitus (Type I), juvenile

diabetes, autoimmune hemolytic anemia, autoimmune

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neutropenia, thrombocytopenia, myasthenia gravis, multiple sclerosis, psoriasis, lichenplanus, graft vs. host disease, acute dermatomyositis, eczema, primary cirrhosis, hepatitis, uveitis, Behcet's disease, acute dermatomyositis, atopic skin disease, pure red cell aplasia, aplastic anemia, amyotrophic lateral sclerosis and nephrotic syndrome.

Preferably the autoimmune condition is glomerulonephritis, insulin-dependent diabetes mellitus (Type I), juvenile diabetes, psoriasis, graft vs. host disease, including transplant rejection, and hepatitis.

Examples of such destructive bone disorders include, but are not limited to, osteoporosis and multiple myeloma-related bone disorder.

Examples of such proliferative conditions include, but are not limited to, acute myelogenous leukemia, chronic myelogenous leukemia, metastatic melanoma, Kaposi's sarcoma, and multiple myeloma.

Examples of such neurodegenerative conditions include, but are not limited to, Alzheimer's disease, Parkinson's disease and Huntington's disease.

Although this invention focuses on the use of the compounds disclosed herein for preventing and treating IL-1, apoptosis, IGIF- and IFN-y-mediated diseases, the compounds of this invention can also be used as inhibitory agents for other cysteine proceases.

The compounds of this invention are also useful as commercial reagents which effectively bind to ICE or other cysteine proteases. As commercial reagents, the compounds of this invention, and their derivatives, may be used to block proteolysis of a target peptide in biochemical or cellular assays for ICE and ICE homologs or may be derivatized to bind to a

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stable resin as a tethered substrate for affinity chromatography applications. These and other uses which characterize commercial cystine protease inhibitors will be evident to those of ordinary skill in the art.

Process of Preparing N-Acylamino Compounds

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The ICE inhibitors of this invention may be synthesized using conventional techniques.

Advantageously, these compounds are conveniently synthesized from readily available starting materials.

The compounds of this invention are among the most readily synthesized ICE inhibitors known.

Previously described ICE inhibitors often contain four or more chiral centers and numerous peptide linkages.

The relative ease with which the compounds of this

invention can be synthesized represents an advantage in the large scale production of these compounds.

For example, compounds of this invention may be prepared using the processes described herein. As can be appreciated by the skilled practitioner, these processes are not the only means by which the compounds described and claimed in this application may be synthesized. Further methods will be evident to those of ordinary skill in the art. Additionally, the various synthetic steps described herein may be performed in an alternate sequence or order to give the desired compounds.

This invention also provides a preferred method for preparing the compounds of this invention. Accordingly, in another embodiment (M) is provided a process for preparing an N-acylamino compound comprising the steps of:

a) mixing a carboxylic acid with an N-

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alloc-protected amino in the presence of an inert solvent, triphenylphoshine, a nucleophilic scavenger, and tetrakis-triphenyl phosphine pailadium(0) at ambient temperature under an inert atmosphere; and

b) adding to the step a) mixture, HOBT and EDC; and optionally comprising the further step of;

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c) hydrolyzing the step b) mixture in the presence of a solution comprising an acid and H2O, wherein the step b) mixture is optionally concentrated, prior to hydrolyzing.

Preferably, the inert solvent is ${\rm CH_2Cl_2},~{\rm DMF},$ or a mixture of ${\rm CH_2Cl_2}$ and DMF.

Preferably, the nucleophilic scavenger is dimedone, morpholine, trimethylsilyl dimethylamine, or 15 dimethyl barbituric acid. More preferably, the nucleophilic scavenger is trimethylsilyl dimethylamine or dimethyl barbituric acid.

Preferably, the solution comprises trifluoroacetic acid in about 1-90% by weight. More preferably, the solution comprises trifluoroacetic acid in about 20-50% by weight.

Alternatively, the solution comprises hydrochloric acid in about 0.1-30% by weight. More preferably, the solution comprises hydrochloric acid in about 0.1-30% by weight.

More preferably, in the above process, the inert solvent is $\mathrm{CH}_2\mathrm{Cl}_2$, DMF, or a mixture of $\mathrm{CH}_2\mathrm{Cl}_2$ and DMF and the nucleophilic scavenger is dimedone, morpholine, trimethylsilyl dimethylamine, or dimethyl barbituric acid.

Most preferably, in the above process the inert solvent is $\mathrm{CH}_2\mathrm{Cl}_2$, DMF, or a mixture of $\mathrm{CH}_2\mathrm{Cl}_2$ and DMF and the nucleophilic scavenger is trimethylsilyl dimethylamine or dimethyl barbituric acid.

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Preferably, the N-acyclamino compound is represented by formula (VIII):

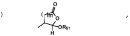
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wherein:

Rl is as defined above in embodiment (A);

R2 is:

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wherein R_{51} is as defined above in embodiment (B);



, or

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Preferably, the N-alloc-protected amine is:

, wherein
$$R_{51}$$
 is as defined above. Aloce—N $$\mathsf{OR}_{\mathfrak{g}}$$

In preferred processes, the substituents are as defined in embodiment (A).

 $\label{eq:local_problem} Alternatively, \ \mbox{the N-acylamino compound is} \\ \ \mbox{represented by formula (VIII), wherein R_1 is as defined}$

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above in embodiment (B) and R_2 is as defined above in embodiment (M).

 $\qquad \qquad \text{Preferably in these alternative} \\ \text{processes, the substituents are as defined above in} \\ \text{embodiment (B).}$

Alternatively, the N-acylamino compound is represented by formula (VIII), wherein R_1 is as defined above in embodiment (C) and R_2 is as defined above in embodiment (M).

 $\label{eq:preferably in these alternative} % \[\begin{array}{c} \text{Preferably in these alternative} \\ \text{processes, the substituents are as defined above in } \\ \text{embodiment (C).} \end{array}$

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Alternatively, the N-acylamino compound is represented by formula (VIII), wherein R_1 is as defined above in embodiment (D) and R_2 is as defined above in embodiment (M).

 $\qquad \qquad \text{Preferably in these alternative} \\ \text{processes, the substituents are as defined above in} \\ \text{embodiment (D).}$

Alternatively, the N-acylamino compound is represented by formula (VIII), wherein R₁ is as defined above in embodiment (E) and R₂ is as defined above in embodiment (M).

Preferably in these alternative
25 processes, the substituents are as defined above in embodiment (E).

Alternatively, the N-acylamino compound is represented by formula (VIII), wherein $\rm R_1$ is as defined above in embodiment (F) and $\rm R_2$ is as defined above in embodiment (M).

 $\label{eq:preferably in these alternative}$ processes, the substituents are as defined above in embodiment (F).

Alternatively, the N-acylamino compound is

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represented by formula (VIII), wherein R_1 is as defined above in embodiment (G) and R_2 is as defined above in embodiment (G).

Preferably in these alternative

processes, the substituents are as defined above in embodiment (G).

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Alternatively, the N-acylamino compound is represented by formula (VIII), wherein $\rm R_1$ is as defined above in embodiment (H) and $\rm R_2$ is as defined above in embodiment (M).

Preferably in these alternative processes, the substituents are as defined above in embodiment (H).

Alternatively, the N-acylamino compound is

15 represented by formula (VIII), wherein R₁ is as defined above in embodiment (I) and R₂ is as defined above in embodiment (M).

 $\label{eq:preferably} \mbox{ In these alternative} \\ \mbox{processes, the substituents are as defined above in embodiment (I).}$

Alternatively, the N-acylamino compound is represented by formula (VIII), wherein $\rm R_1$ is as defined above in embodiment (J) and $\rm R_2$ is as defined above in embodiment (M).

25 Preferably in these alternative processes, the substituents are as defined above in embodiment (J).

Alternatively, the N-acylamine compound is represented by formula (VIII), wherein R_1 is as defined above in embodiment (K) and R_2 is as defined above in embodiment (M).

 $\label{eq:preferably in these alternative} % \[\begin{array}{c} \text{Preferably in these alternative} \\ \text{processes, the substituents are as defined above in embodiment (K).} \end{array}$

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Alternatively, the N-acylamino compound is represented by formula (VIII), wherein ${\bf R}_1$ is as defined above in embodiment (L) and ${\bf R}_2$ is as defined above in embodiment (M).

Preferably in these alternative processes, the substituents are as defined above in embodiment $\{L\}$.

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In order that this invention be more fully understood, the following examples are set forth.

These examples are for the purpose of illustration only and are not to be construed as limiting the scope of the invention in any way.

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Example 1 Inhibition of ICE

We obtained inhibition constants (K_i) and IC_{50} values for compounds of this invention using the three 5 methods described below:

1. Enzyme assay with UV-visible substrate

This assay is run using an Succinyl-Tyr-Val-Ala-Asp-pNitroanilide substrate. Synthesis of analogous substrates is described by L. A. Reiter (Int.

10 J. Peptide Protein Res. 43, 87-96 (1994)). The assay mixture contains:

65 µl buffer (10mM Tris, 1 mM DTT, 0.1% CHAPS @pH 8.1) 10 µl ICE (50 nM final concentration to give a rate of ~lmoD/min)

15 5 µl DMSO/Inhibitor mixture 20 ul 400µM Substrate (80 µM final concentration) 100µl total reaction volume

The visible ICE assay is run in a 96-well microtiter plate. Buffer, ICE and DMSO (if inhibitor

- 20 is present) are added to the wells in the order listed. The components are left to incubate at room temperature for 15 minutes starting at the time that all components are present in all wells. The microtiter plate reader is set to incubate at 37 °C. After the 15 minute
- 25 incubation, substrate is added directly to the wells and the reaction is monitored by following the release of the chromophore (pNA) at 405 - 603 nm at 37 °C for 20 minutes. A linear fit of the data is performed and the rate is calculated in mOD/min. DMSO is only
- 30 present during experiments involving inhibitors, buffer is used to make up the volume to 100 μl in the other experiments.

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2. Enzyme Assay with Fluorescent Substrate

This assay is run essentially according to Thornberry et al. (Nature <u>356</u>: 768-774 (1992)), using substrate <u>17</u> referenced in that article. The substrate

5 is: Acetyl-Tyr-Val-Ala-Asp-amino-4-methylcoumarin (AMC). The following components are mixed:

65 µl buffer(10mM Tris,1mM DTT, 0.1% CHAPS @pH8.1) 10 µl TCE (2 - 10 nM final concentration) 5 µl DMSO/inhibitor solution

0 20 11 150 µM Substrate (30 µM final) 100ul total reaction volume

The assay is run in a 96 well microtiter plate. Buffer and ICE are added to the wells. The components are left to incubate at 37 °C for 15 minutes

- 15 in a temperature-controlled wellplate. After the 15 minute incubation, the reaction is started by adding substrate directly to the wells and the reaction is monitored @37 °C for 30 minutes by following the release of the AMC fluorophore using an excitation
- 20 wavelength for 380 nm and an emission wavelength of 460 nm. A linear fit of the data for each well is performed and a rate is determined in fluorescence units per second.

For determination of enzyme inhibition

25 constants (K₁) or the mode of inhibition (competitive, uncompetitive or noncompetitive), the rate data determined in the enzyme assays at varying inhibitor concentrations are computer-fit to standard enzyme kinetic equations (see I. H. Segel, Enzyme Kinetics, 30 Wiley-Interscience, 1975).

The determination of second order rate constants for irreversible inhibitors was performed by fitting the fluorescence vs time data to the progress equations of Morrison. Morrison, J.F., Mol. Cell.

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<u>Biophys.</u>, 2, pp. 347-368 (1985). Thornberry et al. have published a description of these methods for measurement of rate constants of irreversible inhibitors of ICE. Thornberry, N.A., et al.

- 5 Biochemistry, 33, pp. 3923-3940 (1994). For compounds where no prior complex formation can be observed kinetically, the second order rate constants (k_{inact}) are derived directly from the slope of the linear plots of k_{obs} vs. [I]. For compounds where prior complex
- 10 formation to the enzyme can be detected, the hyperbolic plots of k_{obs} vs. [I] are fit to the equation for saturation kinetics to first generate $K_{\underline{i}}$ and k'. The second order rate constant k_{inact} is then given by $k'/K_{\underline{i}}$.

15 3. PBMC Cell assav

IL-1β Assay with a Mixed Population of Human Peripheral Blood Mononuclear Cells (PBMC) or Enriched Adherent Mononuclear Cells

Processing of pre-IL-1 β by ICE can be

- 20 measured in cell culture using a variety of cell sources. Human PBMC obtained from healthy donors provides a mixed population of lymphocyte subtypes and mononuclear cells that produce a spectrum of interleukins and cytokines in response to many classes
- 25 of physiological stimulators. Adherent mononuclear cells from PBMC provides an enriched source of normal monocytes for selective studies of cytokine production by activated cells.

Experimental Procedure:

An initial dilution series of test compound in DMSO or ethanol is prepared, with a subsequent dilution into RPMI-10% FBS media (containing 2 mM L-glutamine, 10 mM HEPES, 50 U and 50 ug/ml pen/strep)

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respectively to yield drugs at 4x the final test concentration containing 0.4% DMSO or 0.4% ethanol. The final concentration of DMSO is 0.1% for all drug dilutions. A concentration titration which brackets 5 the apparent $K_{\rm i}$ for a test compound determined in an ICE inhibition assay is generally used for the primary compound screen.

Generally 5-6 compound dilutions are tested and the cellular component of the assay is performed in 10 duplicate, with duplicate ELISA determinations on each cell culture supernatant.

PBMC Isolation and IL-1 Assay:

Buffy coat cells isolated from one pint human blood (yielding 40-45 ml final volume plasma plus cells) are diluted with media to 80 ml and LeuxoPREP separation tubes (Becton Dickinson) are each overlaid with 10 ml of cell suspension. After 15 min centrifugation at 1500-1800 xg, the plasma/media layer is aspirated and then the mononuclear cell layer is collected with a Pasteur pipette and transferred to a 15 ml conical centrifuge tube (Corning). Media 1s added to bring the volume to 15 ml, gently mix the cells by inversion and centrifuge at 300 xg for 15 min. Resuspend the PBMC pellet in a small volume of media, 25 count cells and adjust to 6 x 10⁶ cells/ml.

For the cellular assay, 1.0 ml of the cell suspension is added to each well of a 24-well flat bottom tissue culture plate (Corning), 0.5 ml test compound dilution and 0.5 ml LFS solution (Sigma 30 #L-3012; 20 ng/ml solution prepared in complete RPMI media; final LFS concentration 5 ng/ml). The 0.5 ml additions of test compound and LFS are usually sufficient to mix the contents of the wells. Three control mixtures are run per experiment, with either

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LPS alone, solvent vehicle control, and/or additional media to adjust the final culture volume to 2.0 ml. The cell cultures are incubated for 16-18 hr at 37 $^{\circ}$ C in the presence of 5% CO₂.

5 At the end of the incubation period, cells are harvested and transferred to 15 ml conical centrifuge tubes. After centrifugation for 10 min at 200 xg, supernatants are harvested and transferred to 1.5 ml Eppendorf tubes. It may be noted that the cell pellet may be utilized for a biochemical evaluation of pre-IL-1β and/or mature IL-1β content in cytosol extracts by western blotting or ELISA with pre-IL-1β specific antisera.

Isolation of Adherent Mononuclear cells:

above. Media (1.0 ml) is first added to wells followed by 0.5 ml of the PBMC suspension. After a one hour incubation, plates are gently shaken and nonadherent cells aspirated from each well. Wells are then gently washed three times with 1.0 ml of media and final resuspended in 1.0 ml media. The enrichment for adherent cells generally yields 2.5-3.0 x 10⁵ cells per well. The addition of test compounds, LPS, cell incubation conditions and processing of supernatants

ELISA:

We have used Quantikine kits (R&D Systems) for measurement of mature IL-1β. Assays are performed according to the manufacturer's directions. Mature 30 IL-1β levels of about 1-3 ng/ml in both PBMC and adherent mononuclear cell positive controls are observed. ELISA assays are performed on 1:5, 1:10 and 1:20 dilutions of supernatants from LFS-positive

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controls to select the optimal dilution for supernatants in the test panel.

The inhibitory potency of the compounds can be represented by an IC_{5C} value, which is the concentration of inhibitor at which 50% of mature IL-1

5 concentration of inhibitor at which 50% of mature IL-1 β is detected in the supernatant as compared to the positive controls.

 $\label{thm:continuous} The \ skilled \ practitioner \ realizes \ that \ values \\ obtained \ in \ cell \ assays, \ such \ as \ those \ described$

10 herein, can depend on multiple factors, such as cell type, cell source, growth conditions and the like.

Example 2

Pharmacokinetic Studies in the Mouse

- Peptidyl ICE inhibitors are cleared rapidly with clearance rates greater than 100 µ/min/kg. Compounds with lower clearance rates have improved pharmacokinetic properties relative to peptidyl ICE inhibitors.
- 20 We obtained the rate of clearance in the mouse (μ/min/kg) for several compounds of this invention using the method described below:

Sample Preparation and Dosing

Compounds were dissolved in sterile TRIS

25 solution (0.02M or 0.05M) at a concentration of
 2.5mg/ml. Where necessary to ensure a complete
 solution, the sample was first dissolved in a minimum

solution, the sample was first dissolved in a minimum of dimethylacetamide (maximum of 5° of total solution volume) then diluted with the TRIS solution.

30 The drug solution was administered to CD-1 mice (Charles River Laboratories - 26-31q) via the tail

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vein at a dose volume of 10ml/kg giving a drug dose of 25mg/kg.

Mice were dosed in groups of 5 for each timepoint (generally from 2 minutes to 2 hours) then at 5 the appropriate time the animals were anaesthetised with halothane and the blood collected into individual heparinized tubes by jugular severance. The blood samples were cooled to 0 °C then the plasma separated and stored at -20 °C until assayed.

10 Bioassay

Drug concentration in the plasma samples were determined by HPLC analysis with UV or MS (ESP)

detection. Reverse phase chromatography was employed using a variety of bonded phases from C1 to C18 with 15 eluents composed of aqueous buffer/acetonitrile

mixtures run under isocratic conditions.

Quantitation was by external standard methods with calibration curves constructed by spiking plasma with drug solutions to give concentrations in the range 20 of 0.5 to $50\mu g/ml$.

Prior to analysis the plasma samples were deproteinated by the addition of acetonitrile, methanol, trichloroacetic acid or perchloric acid followed by centrifugation at 10,000g for 10 minutes.

25 Sample volumes of $20\mu l$ to $50\mu l$ were injected for analysis.

Compound 214e

Dosing and sampling

The drug was dissolved in sterile 0.02M Tris
to give a 2.5mg/ml solution which was administered to
11 groups of 5 male CD-1 mice via the tail vein at a
dose of 25mg/kg. At each of the following timepoints:
2, 5, 10, 15, 20, 30, 45, 60, 90 and 120 minutes a

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group of animals was anaesthetised and the blood collected into heparinized tubes. After separation the plasma was stored at $-20\,^{\circ}\mathrm{C}$ until assayed.

Assay

Aliquots of plasma (150µ1) were treated with 5% perchloric acid (5µ1) then mixed by vortexing and allowed to stand for 90 minutes prior to centrifugation. The resulting supernatant was separated and 20µ1 was injected for HPLC analysis.

10 HPLC Conditions

Column 100 x 4.6mm Kromasil KR 100 5C4 Mobile Phase 0.1m Tris pH7.5 86%

Acetonitrile 14%

Flowrate 1ml/min

15 Detection UV at 210nm Retention Time 3.4 mins

The results of the analysis indicated a decrease in the mean plasma level of the drug from \sim 70 μ g/ml at 2 minutes to < 2 μ g/ml at 90 and 120 minutes.

20 Compound 217e

Dosing and sampling

The drug was dissolved in sterile 0.02M Tris to give a 2.5 mg/ml solution which was administered to 11 groups of 5 male CD-1 mice via the tail vein at a

25 dose of 25mg/kg. At each of the following timepoints: 2, 5, 10, 15, 20, 30, 45, 60, 90 and 120 minutes a group of animals was anaesthetised and the blood collected into heparinized tubes. After separation the plasma was stored at -20 °C until assayed.

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Assav

Aliquots of plasma (100µ1) were diluted with acetonitrile (100µ1) then mixed by vortexing for 20 seconds before centrifugation for 10 minutes. The 5 resulting supernatant was separated and 20µ1 was injected for HPLC analysis.

HPLC Conditions

Column 150 x 4.6mm Zorbax SBC8 Mobile Phase 0.05M Phosphate 72%

10 buffer ph7.1

Acetonitrile 28%

Flowrate 1.4ml/min
Detection UV at 210nm

Retention Time 3.0 and 3.6 mins (diasteromers)

The results of the analysis indicated a decrease in mean plasma concentrations from ~ 55ug/ml at 2 minutes to < 0.2ug/ml at 60-120 minutes.</p>

Example 3

Peptidyl ICE inhibitors are cleared rapidly
with clearance rates greater than 80 ml/min/kg.
Compounds with lower clearance rates have improved
pharmacokinetic properties relative to peptidyl ICE
inhibitors.

We obtained the rate of clearance in the rat 25 (ml/min/kg) for several compounds of this invention using the method described below:

In vivo Rat Clearance Assay

Cannulations of the jugular and carotid vessels of rats under anesthesia were performed one day 30 prior to the pharmacokinetic study. M.J. Free, R.A.

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Jaffee; 'Cannulation techniques for the collection blood and other bodily fluids'; in: Animal Models; p. 480-495; N.J. Alexander, Ed.; Academic Press; (1978). Drug (10mg/mL) was administered via the

- 5 Jugular vein in a vehicle usually consisting of: propylene glycol/saline, containing 100mM sodium bicarbonate in a 1:1 ratio. Animals were dosed with 10-20 mg drug/kg and blood samples were drawn at 0, 2, 5, 7, 10, 15, 20, 30, 60, and 90 minutes from an
- 10 indwelling carotid catheter. The blood was centrifuged to plasma and stored at -20 °C until analysis. Pharmacokinetic analysis of data was performed by nonlinear regression using standard software such as RStrip (MicroMath Software, UT) and/or Pcnonlin (SCI
- 15 Software, NC) to obtain clearance values.

Analytical:

Rat plasma was extracted with an equal volume of acetonitrile (containing 0.1% TFA). Samples were then centrifuged at approximately 1,000 x g and the

20 supernatant analyzed by gradient HPLC. A typical assay procedure is described below.

 $200~\mu L$ of plasma was precipitated with 200 μL of 0.1% trifluoroacetic acid (TFA) in acetonitrile and 10 μL of a 50% aqueous zinc chloride solution, vortexed

25 then centrifuged at ~1000 x g and the supernatancellected and analyzed by HPLC.

HPLC procedure:

Column: Zorbax SB-CN (4.6 x 156 mm) (5u

30 Column temperature: 50 °C Flow rate: 1.0 mL/min Injection volume: 75 μL .

Mobile phase: $A=0.1^{\circ}$ TFA in water and B=100

acetonitrile

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Gradient employed: 100% A to 30% A in 15.5 min 0% A at 16 min

100% A at 19.2 min

Wavelength: 214 nm

A standard curve was run at 20, 10, 5, 2 and 1 $\mu q/mL$ concentrations.

Example 4

Whole Blood Assay for IL-18 Production

 $\label{eq:weak_compounds} \mbox{We obtained IC_{50} values for several compounds} \\ \mbox{10} \mbox{ of this invention using the method described below:}$

Purpose:

The whole blood assay is a simple method for measuring the production of IL-lb (or other cytokines) and the activity of potential inhibitors. The

- 15 complexity of this assay system, with its full complement of lymphoid and inflammatory cell types, spectrum of plasma proteins and red blood cells is an ideal <u>in vitro</u> representation of human <u>in vivo</u> physiologic conditions.
- 20 Materials:

Pyrogen-free syringes (~ 30 cc)

Pyrogen-free sterile vacuum tubes containing
lyophilized Na₂EDTA (4.5 mg/10 ml tube)

Human whole blood sample (~ 30-50 cc)

25 1.5 ml eppendorf tubes Test compound stock solutions (~ 25mM in DMSO or other solvent) Endotoxin-free sodium chloride solution (0.9%) and HBSS

Lipopolysaccharide (Sigma; Cat.# L-3012) stock solution

30 at lmg/ml in HBSS

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IL-1 β ELISA Kit (R & D Systems; Cat # DLB50) TNF α ELISA Kit (R & D Systems; Cat # DTA50) Water bath or incubator

Whole Blood Assav Experimental Procedure:

Set incubator or water bath at 30 °C.

Aliquot 0.25ml of blood into 1.5 ml eppendorf tubes.

Note: be sure to invert the whole blood sample tubes after every two aliquots. Differences in replicates may result if the cells sediment and are not uniformly

10 suspended. Use of a positive displacement pipette will also minimize differences between replicate aliquots. Prepare drug dilutions in sterile pyrogen-

free saline by serial dilution. A dilution series which brackets the apparent $K_{\mathbf{i}}$ for a test compound

- 15 determined in an ICE inhibition assay is generally used for the primary compound screen. For extremely hydrophobic compounds, we have prepared compound dilutions in fresh plasma obtained from the same blood donor or in PBS-containing 5% DMSO to enhance
- 20 solubility.

Add 25 µl test compound dilution or vehicle control and gently mix the sample. Then add 5.0 µl LPS solution (250 ng/ml stocked prepared fresh: 5.0 ng/ml final concentration LPS), and mix again. Incubate the 25 tubes at 30 °C in a water bath for 16-18 hr with occasional mixing. Alternatively, the tubes can be placed in a rotator set at 4 rpm for the same

- occasional mixing. Alternatively, the tubes can be placed in a rotator set at 4 rpm for the same incubation period. This assay should be set up in duplicate or triplicate with the following controls:
- 30 negative control- no LPS; positive control- no test inhibitor; vehicle control- the highest concentration of DMSO or compound solvent used in the experiment.

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Additional saline is added to all control tubes to normalize volumes for both control and experimental whole blood test samples

After the incubation period, whole blood .

5 samples are centrifuged for 10 minutes at ~ 2000 rpm in the microfuge, plasma is transferred to a fresh microfuge tube and centrifuged at 1000 x g to pellet residual platelets if necessary. Plasma samples may be stored frozen at -70 °C prior to assay for cytokine 10 levels by ELISA.

ELISA:

We have used R & D Systems (614 McKinley Place N.E. Minneapolis, MN 55413) Quantikine kits for measurement of IL-1β and TNF-α. The assays are 15 performed according to the manufacturer's directions. We have observed IL-1β levels of ~ 1-5 ng/ml in positive controls among a range of individuals. A 1:200 dilution of plasma for all samples has been sufficient in our experiments for ELISA results to fall 20 on the linear range of the ELISA standard curves. It may be necessary to optimize standard dilutions if you observe differences in the whole blood assay. Nerad, J.L. et al., J. Leukocyte Biol., 52, pp. 687-692 (1992).

25 Example_5

Inhibition of ICE homologs

Isolation of ICE homologs
 Expression of TX in insect cells using a baculovirus expression system. We have subcloned Tx cDNA (Faucheu 30 et al., supra 1995) into a modified pVL1393 transfer

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vector, co-transfected the resultant plasmid
(pVL1393/TX) into insect cells with viral DNA and
identified the recombinant baculovirus. After the
generation of high titer recombinant virus stock, the
5 medium was examined for TX activity using the visible
ICE assay. Typically, infection of Spodoptera
frugiperda (Sf9) insect cells at an MOI of 5 with
recombinant virus stock resulted in a maximum
expression after 48 hours of 4.7µg/ml. ICE was used as
10 a standard in the assay.

Amino terminal T7 tagged versions of ICE or TX were also expressed. Designed originally to assist the identification and purification of the recombinant proteins, the various constructs have also allowed

15 examination of different levels of expression and of the relative levels of apoptosis experienced by the different homologs. Apoptosis in the infected Sf9 cells (examined using a Trypan Blue exclusion assay) was increased in the lines expressing ICE or TX

20 relative to cells infected with the viral DNA alone.

Expression and purification of N-terminally (His)6tagged CPP32 in E. coli. A cDNA encoding a CPP32
(Fernandes-Alnemri et al, supra 1994) polypeptide
starting at Ser (29) was PCR amplified with primers
that add in frame XhoI sites to both the 5' and 3' ends
of the cDNA and the resulting XhoI fragment ligated
into a Xho I-cut pET-15b expression vector to create an
in frame fusion with (his)6 tag at the n-terminus of
the fusion protein. The predicted recombinant protein
starts with the amino acid sequence of
MGSSHHHHHHHSSGLVPRGSHMLE, where LVPRGS represents a
thrombin cleavage site, followed by CPP32 starting at

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Ser (29). E. coli BL21(DE3) carrying the plasmid were grown to log phase at 30 °C and were then induced with 0.8 mM IPTG. Cells were harvested two hours after IPTG addition. Lysates were prepared and soluble proteins 5 were purified by Ni-agarose chromatography. All of the expressed CPP32 protein was in the processed form. N-terminal sequencing analysis indicated that the processing occurred at the authentic site between Asp (175) and Ser (176). Approximately 50 µg of CPP32

10 protein from 200 ml culture. As determined by active site titration, the purified proteins were fully active. The protease preparation were also very active in vitro in cleaving PARP as well as the synthetic DEVD-AMC substrate (Nicholson et al, supra 1995).

15 2. Inhibition of ICE homologs

The selectivity of a panel of reversible inhibitors for ICE homologs is depicted in Table 1. ICE enzyme assays were performed according to Wilson et al (<u>supra 1994</u>) using a YVAD-AMC substrate (Thornberry et al, supra

- 20 1992). Assay of TX activity was performed using the ICE substrate under identical conditions to ICE. Assay of CPP32 was performed using a DEVD-AMC substrate (Nicholson et al., <u>supra</u> 1995). In general, there is low selectivity between ICE and TX for a wice range of
- 25 scaffolds. None of the synthetic ICE compounds tested are effective inhibitors of CPP32. Assay of the reversible compounds at the highest concentration (1 um) revealed no inhibition.

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Table 1

	Compound	K _i ICE (nM)	K _i TX (nM)	K _i CPP32 (nM)
	214e	7.5	7.0 ± 1.1	> 1000
	135a	90	55 ± 9	>1000
5	125b	60	57 ± 13	> 1000
	137	40	40 ± 7	> 1000

Second-order rate constants for inactivation of ICE and ICE homologs with selected irreversible inhibitors are presented below (Table 2). The

10 irreversible compounds studied are broad spectrum inhibitors of ICE and its homologs. Some selectivity, however, is observed with the irreversible compounds comparing inhibition of ICE and CPP32.

Table 2

15	Compound	k _{inact} (ICE) M ⁻¹ s ⁻¹	k _{inact} (TX) k _{inact} (CPP32) M ⁻¹ s ⁻¹
	138	120,000	150,000	550,000
	217d	475,000	250,000	150,000
	108a	100,000	25,000	nd

Example 6

20 Inhibition of apoptosis

Fas-Induced Apoptosis in U937 cells. Compounds were evaluated for their ability to block anti-Fas-induced apoptosis. In a preliminary experiment using RT-PCR, 25 we detected mRNA encoding ICE, TX, ICH-1, CPP32 and

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CMH-1 in unstimulated U937 cells. We used this cell line for apoptosis studies. U937 cells were seeded in culture at 1 x 10⁵ cells/ml and grown to ~5 x 10⁶ cells/ml. For apoptosis experiments, 2 x 10⁶ cells were plated in 24-well tissue culture plates in 1 ml RPMI-1640-10% FBS and stimulated with 100 ng/ml anti-Fas antigen antibody (Medical and Biological Laboratories, Ltd.). After a 24 hr incubation at 37 °C, the percentage of apoptotic cells was determined by 10 FACS analysis using ApoTag reagents.

All compounds were tested initially at 20 μ M and titrations were performed with active compounds to determine IC $_{50}$ values. Inhibition of apoptosis (> 75% at 20 μ M) was observed for 108a, 136, and 138.

15 An IC50 of 0.8 μ M was determined for 217e compared to no inhibition of anti-Fas-induced apoptosis by 214e at 20 μ M.

Example 7

In vivo acute assay for efficacy as anti-inflammatory agent

LPS-Induced IL-18 Production.

20

Efficacy of 214e and 217e was evaluated in CD1 mice (n=6 per condition) challenged with LPS (20 mg/kg IP). The test compounds were prepared in clive 25 oil:DMSO:ethanol (90:5:5) and administered by IP injection one hour after LPS. Blood was collected seven hours after LPS challenge. Serum IL-1β levels were measure by ELISA. Results in Fig. 6 show a dose dependent inhibition of IL-1β secretion by 214e, with

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an ED $_{50}$ of approximately 15 mg/kg. Similar results were obtained in a second experiment. A significant inhibition of IL-1 β secretion was also observed in 217e treated mice (Fig. 7). However, a clear dose response 5 was not apparent.

Compounds 214e and 217e (50 mg/kg) were also administered by oral gavage to assess absorption. Results in Fig. 8 show that 214e, but not 217e when administered orally inhibited IL-1ß secretion, suggesting potential for oral efficacy of ICE inhibitors as anti-inflammatory agents.

The efficacy of analogs of 214e were also evaluated in LPS challenged mice after IP administration (Fig. 9) and PO administration

Table 3 % Inhibition of IL- β production by analogs of 214e in LPs-chellenged mice after PO and IP administration (50 mg/kg).

Table 3

Compound	PO% Inhibition	IP% Inhibition
214e	75	78
265	27	30
416	52	39
434	80	74
438	13	40
442	10	Ü
2002	-	78

25

20

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Table 4

Comparison of 214e Prodrugs for Efficacy in LPS Challenged Mice: Time Course Inhibition of IL-1 β Production

Time of Compound Administration (relative to time of LPS challenge, PO, 50 mg/kg

5	Compound	-2 hr	-1 hr	0 hr	+1 hr
	214e	39* 43* -*	-* 44* -*	80* 48* -*	55% 75* 11* 47*
	304a	30	33	68	37
	2100e	49	54	94	66
	2100a	8	71	67	58
10	213e	0	48	41	89
	302	0	27	21	26
	2100c	0	0	85	40
ĺ	2100d	42	35	52	26
	2100b	0	0	47	26
15	2001	~63 64*	~62 62*	~57 58*	~54 55*

^{*} Values obtained in subsequent assays

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Example 8

Measurement of blood levels of prodrugs of 214e.

Mice were administered a p.o. dose of compounds 302 and 304a (50 mg/kg) prepared in 0.5 % 5 carboxymethylcellulose. Blood samples were collected at 1 and 7 hours after dosing. Serum was extracted by precipitation with an equal volume of acetonitrile containing 2 % formic acid followed by centrifugation. The supernatant was analyzed by liquid chromatography-10 mass spectrometry (ESI-MS) with a detection level of 0.03 to 3 µg/ml. Compounds 302 and 304a showed detectable blood levels when administered orally, 214e itself shows no blood levels above 0.10 µg/mL when administered orally. Compounds 302 and 304a are

Example 9

We obtained the following data (see Tables 5 and 6) for compounds of this invention using the 20 methods described in Examples 1-8. The structures of the compounds of Example 9 are shown in Example 10-17.

Table 5

25

(see Fig. 11).

Compoun		UV- isible 1 (nM)	Cell PBMC avg. IC50 (nM)	Whole human blood IC50 (nM)	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
47b	1	27	1800	<600	338	
47a		19	2600	5100	79	32
135a		90	2800	5000	>100	

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	Compound	UV- Visible Ki (nM)	Cell PBMC avg. IC50 (nM)	Whole human blood IC50 (nM)	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
	135b	320	1600	1700		
	125b	60	800	4500		
	108b	400	25000	1		>100
	137	40	1700	14000		
5	139	350	2000	1		
	213e	130	900	600 400*		
	214c	1200	5000			
	214e	7.5	1600	1300	23	12
	217c		1700	7000	70	
10	217e		175	2000	>50	-
	220b	600	2125			
	223b	99	5000		>100	
	223e	1.6	3000	>20000	89	
	226e	15	1100	1800	109	
15	227e	7	234	550		
	230e		325	300	67	
	232e	1100	4500		22	26
	235e	510	4750		36	
	238e	500	4250		-	
20	246	12	950	10000	31	
	257	13	11000 6600*			
	265	47	4300	1400	23	20
	281	50	600 2500*			
	302	4500	>20000	>20000		
25	304a	200	1,400	2400 14000*		
	307a	55	14500	16000		
	307b	165		14000		
	404	2.9	1650 1800*	1100	64	24
	405	6.5	1700	2100		
30	406	4	1650	2300		
	407	0.4	540	1700		
	408	0.5	1100	1000	41	23

	-	UV-	Cell PBMC	Whole	.Clearance	Clearance
	Compound		avq.	blood	Mouse,	ml/min/kg
	compound	Ki (nM)	IC50	IC50	i.v.	
		112 (121)	(nM)	(nM)	ml/min/kg	1
	409	3.7	2500			
	410	17	2000	2800	32	20
	411	0.9	540	1900		
	412	1.3	580	700		25
			660*	1000*		
5	413	750	6200			
	415	2.5	990	450	26	18
	-		1000*	3500*		
	416	12	1200	3400		47
	417	8	2000	6000	33	22
	418	2.2	1050 2200*	7800 1800*	13	5.9
10	419	280	>8000	1000		
10	420	1200	8000			
	420	1200	>8000*			
	421	200	4300 4600*			
	422	50	2200	1200		
	423	10	2100 1800*	1500		4.5
15	424	45	2500	4000		
10	425	0.8	650	650		
	423	0.0	700*	030		
	426	90	4500 2500*			
	427	180	4500			36
	428	280				
20	429	7000				
	430	60 ,	>8000			
	431	8	>8000	8000		
	432	1.6	560	2000		
	433	2.9	1000 1100*	1100		
25	434	4.9	1600	1800		20
			1200*	1300*		2.0
	435	8	4400			
	436	7.5 .	2700			
	437	12	1800	5000		

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	UV-	Cell PBMC	Whole	Clearance	Clearance Rat, i.v.
Compo	und Visible	avg.	blood	Mouse,	ml/min/ko
1	Ki (nM)	IC50	IC50	ml/min/kg	
		(nM)	(nM)	MIT/MITH/ AG	!
438	28	1000	700		22
-			2900+	·	-
439	3.7	2800	3200	!	I I
440	2.3	5000	3400*	-	
441		2500	2000	·	
			4500		
442		900	2000		54
443		2800	1500	<u> </u>	
444		3500	3500	1	
445			4000		
446	62		3000		
447	5.8	2500	1500		
448	130		4000		
449	12	1500	3200		
			13000*		
450	5	800	2200	18	12
			1700*		
451	4	1800	1500		
			9000*		
452	4.5	600	650		27.3
450	0.65	800*	1600*	<u> </u>	
453	0.65	1300	1900 1600*		
454	45	2500	1000-		
455	1.2	400	2800		
433	1.2	400	2600*		54
456	4.5	600	600		12.7
		1300*	1400*		
457	6.2	2000	3500		
458	20	2900			
459	5 ,	1800			
460	115	400	2400		
461	47				
462	40				1 months
463	14	2400			
	:	2800*			
464	2.5	1000	>1000		
			2500*		
465	3	1000	800		

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	Compound	UV- Visible Ki (nM)	PBMC avg. IC50 (nM)	Whole human blood IC50 (nM)	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
	466	0.8	1400	600	-	
	467	11	1900			
	468	4.5	850	2500	·	
	470	5	500	360		63
	4,0	, ,	300	500*		63
5	471	1	750	400		17
	472	140				
	473	1	1000	400 450*		
	474	85				
	475	5.5	690 650*	400 350*	31	21
10	476	7	1600	2500		
	477	60				
	478	380				
	479	15	900	700 2400*		
	480	25	2300			
15	481	1.2	390 930*	600 500*		34
	482	<0.2	340	380 260*		
	483	1.7	900	700		
	484	2	1550 1400*	5000		15
	485	2	900	900		
20	486	2.3	480 570*	500		37
	487	2.4	650 950*	500 400*		20
	488	1.5	940	750		
	489	6	2250 1700*	15000		
	490	4.3	980 1000*	700 1900*		
25	491	. 5	2500			
	493	25	1200	800 850*		

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		UV-	Cell PBMC	Whole human	Clearance Mouse,	Clearance Rat, i.v.
	Compound	Ki (nM)	avg. IC50 (nM)	IC50 (nM)	i.v. ml/min/kg	ml/mir/kg
	494	15	1350 1500*	7000		
	495	43				
	496	16	1550 1600*	6000		
	497	3.5	740	350 700*		-
5	498	1.5	560	500 400*		
	499	3.5	1200 800*	9000		
	605a	90	2600	>20000		
	605b	45	10000		97	
	605c	615	4500		37	
10	605d	95	5100	16000 5100*	33	
	605e	29	2250	>10000		24
	605f	475	12500			
	605g	165	22500			
	605h	460	>25000			
15	605i	680	>20000			
	605j	110	8750		71	
	605m	650	20000	1		
	605n	12	2100	>20000	28	
	6050	72		18000		
20	605p	125	3200	>20000		
	605q	1000				
	605s	150	6000			
	605t	33				
	609a	114	>30000			
25	609b	27	>20000			
	619	300				
	620	35	1000	19000		
	621	7.2	1300	>20000		
	622	35	1300	>20000		
30	623	9 .				
	624	300				

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	Compound	UV- Visible Ki (nM)	Cell PBMC avg. IC50 (nM)	Whole human blood IC50 (nM)	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
	625	105		1		
	626	260		1		
	627	43	3250	8000		
	628	36	2750	>20000		
5	629	230				
	630	270				
	631	805				
	632	148				
	633	92	5750	20000		
10	634	1400		1		
	635	55	1900 3400*	4000		
	605v	1100	>30000		!	
	2201	9	2000 3700*	350C		60
	2100e	250	800	600		
15	2100a	100	1100	850	1	
	2002	4	810 860*	70 1400*		32
	2100d	>100000	>20000	>20000	-	
	2100c	7400	>20000	>20000		
	2100b	8000	>20000	>20000		
20	2001	135	1800	3500		
	1027	4000	>20000	>20000		60
	1015	40	2500	1700		23

Table 6

	Compound	Fluorescent Assay k _{inact} M ⁻¹ s ⁻¹	PBMC	Whole human blood IC50 (nM)	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
25	108a	1×10 ⁵	17500			
	136	5.4x10 ⁵	870	2800	93	

Compound	Fluorescent Assay ^k inact M ⁻¹ s ⁻¹	Cell PBMC avg. IC50 (nM)	Whole human blood IC50 (nM)	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
138	1.2x10 ⁵	900	2900	116	
217d	4.7x10 ⁵	340	4000		
280	4×10 ⁵	650	>1000		187
283	1x10 ⁵	<200	450		104
284	3.5x10 ⁵	470	550	77	100
285	4.3x10 ⁵	810	1000	130	50

* Values obtained upon reassay.

5

Example 10

 $\mbox{ Compound 139 was synthesized by a method} \\ 10 \mbox{ similar to the method used to synthesize 47a.}$

Compounds 136 and 138 were synthesized by a method similar to the method used to synthesize 57b.

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Compounds 135a, 135b, and 137 were synthesized by a method similar to the method used to synthesize 69a.

Compounds 813e, 814c, 814e, 817c, 817d, 817e, 820b, 823b, 823e, 826e, 827e, 830e, 832e, 835e, 838e, 10 846, 857, 865, 902, 904a, 907a, 907b, 1004-1013, 1015-

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1045, 1046-1068, 1070-1091, and 1093-1099 were synthesized by methods similar to those used to synthesize compound 264 and the corresponding compounds in Examples 10 and 11.

- 5 Compounds 47a, 47b, 108a, 108b, 125b, 213e, 214c, 217c, 217d, 217e, 220b, 223b, 223e, 226e, 227e, 230e, 232e, 235e, 238e, 246, 257, 264, 265, 280-287, 302, 304a, 307a, and 307b were synthesized as described below.
- 10 H. N-(N-Acetyl-tyrosinyl-valinyl-pipecolyi)-3-amino-4-oxobutanoic acid.
 - Step A. N=(N-tert-Butoxycarbonylpipecolyl)-4amino-5-benzyloxy-2-oxotetrahydrofuran.

Reaction of N-tert-butoxycarbonylpipecolic

acid (460 mg, 2.0 mmol) and N-allyloxycarbonyl-4-amino5-benzyloxy-2-oxotetrahydrofuran (530 mg, 1.82 mmol)
was carried out by a method analogous to that reported
by Chapman (Bicorg. & Med. Chem. Lett. 2, pp. 613-618,
(1992)) to give 654 mg of the title compound.

- 25 Step B. N-Pipecolyl-4-amino-5-benzyloxy-2oxotetrahydrofuran.

N-(N-tert-Butoxycarbonylpipecoly1)-4-amino-5benzyloxy-2-oxo-tetrahydrofuran (654 mg) was dissolved in 15 ml of 25% trifluoroacetic acid in dichloromethane

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and stirred at room temperature. The mixture was concentrated to give a gummy residue. The residue was dissolved in dichloromethane and washed with 10% sodium bicarbonate. The organic layer was dried over

5 anhydrous sodium sulfate, filtered, and concentrated to give 422 mg of the title compound as a beige solid.

¹H NMR (500 MHz, CDCl₃) δ 7.38 (m, 5H), 7.15 (d, 1H), 5.55 (d, 1H), 4.95-4.8 (m, 1H), 4.78 (m, 1H), 4.65 (d, 1H), 4.45 (m, 1H), 3.2 (m, 0.5H), 3.05 (m, 10 0.5H), 2.95 (m, 0.5H), 2.85 (m, 0.5H), 2.65 (m, 1H), 2.55-2.38 (m, 1H), 1.95 (m, 1H), 1.8 (m, 1H), 1.6 (m, 2H), 1.38 (m, 2H).

Step C. N-(N-Acetyl-tyrosinyl-valinylpipecolyl)-4-amino-5-benzyloxy-2-oxotetrahydrofuran.

(SiO₂) eluting with 94:6:1

1.5

N-Acetyl-tyrosinyl-valine (464 mg, 1.44 mmol) and N-Pipecolyl-4-amino-5-benzyloxy-2- oxotetrahydrofuran (412 mg, 1.3 mmol) were dissolved in 5 ml each of dimethylformamide and dichloromethane and 20 cooled to 0°C. To the cooled solution was added 1-hydroxybenzotriazole (HOBT; 210 mg, 1.56 mmol) followed by the addition of 1-(3-dimethylaminopropyl)-3-ethyl carbodiimide hydrochloride (EDC; 326 mg, 1.7 mmol). After stirring for 18 hours, the mixture was 25 diluted with ethyl acetate and washed with water, 10. sodium hydrogen sulfate, 10% sodium bicarbonate, and water. The organic layer was concentrated to give a crude solid that was purified by flash chromatography

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(dichloromethane:isopropanol:pyridine) to give 370 mg of the title compound.

1H NMR (500 MHz, CD₃OD (existing as
diastereomers as well as rotamers)) δ 7.35 (m, 5H),
5 7.05 (m, 2H), 6.68 (m, 2H), 5.65 & 5.25 (m, 1H), 4.93.95 (m, 8H), 3.4-2.6 (m, 4H), 2.5-2.1 (m, 1H), 1.98
(s, 1H), 1.9 (s, 1H), 1.85 (s, 1H), 1.8-1.6 (m, 2H),

Step D. N-(N-Acetyl-tyrosinyl-valinyl-

1.55-1.3 (m, 4H), 0.95-0.85 (m, 6H).

10

20

pipecolyl)-3-amino-4-oxobutanoic acid.

To a solution of 100 mg of N-(N-Acetyl-tyrosinyl-valinyl-pipecolyl)-4-amino-5-benzyloxy-2-oxotetrahydrofuran in 10 ml of methanol was added 60 mg of Pd(OH)2 on carbon and the mixture placed under an atmosphere of hydrogen via a balloon. The mixture was filtered through Celite and concentrated providing a white solid. This crude solid was dissolved in 2 ml of methanol and triturated with diethyl ether affording 26 mg of the title compound.

 ^{1}H NMR (500 MHz, CD₃OD(existing as diastereomers as well as rotamers)) δ 7.1 (m, 2H), 6.7 (m, 2H), 5.2 (br. m, 1H), 4.8-3.6 (m, 6H), 3.2-2.5 (m, 4H), 2.5-2.1 (m, 1H), 1.95 (three s, 3H), 1.9-1.3 (m, 6H), 1.1-0.7 (m, 6H).

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K. N-[N-Acetyl-tyrosinyl-valinyl-(4-

benzyloxy)prolinyl1-3-amino-4-oxobutanoic acid.

Step A. N-(N-Allyloxycarbonyl-4-

benzyloxyprolinyl)-3-amino-4-oxobutanoic

5 acid tert-butyl ester semicarbazone.

The title compound was prepared by the reaction of N-allyloxycarbonyl-4-benzyloxyproline and 3-amino-4-oxobutanoic acid tert-butyl ester semicarbazone (T.L. Graybill et. al., Abstracts of

10 papers, 206th National Meeting of the American Chemical Society, Abstract MEDI-235. Chicago, 1L. (1993)) under similar peptide coupling conditions as reported above (compound H; Step C).

¹H NMR (500 MHz, CDC1₃) δ 9.05 (br. s, 1H),

15 7.85 (br. m, 1H), 7.4-7.2 (m, 5H), 7.15 (br. s, 1H), 6.55 (br. s, 1H), 5.9 (m, 1H), 5.1-4.9 (br. m, 2H), 4.65-4.4 (m, 4H), 4.2 (br. m, 1H), 3.75-3.5 (m, 2H), 2.75-2.55 (m, 2H), 2.5 (br. m, 1H), 2.25 (br. m, 1H) 1.4 (s. 9H).

20 Step B. N-(N-Acetyl-tyrosinyl-valinyl-(4-benzyloxyprolinyl))-3-amino-4oxobutanoic acid tert-butyl ester semicarbazone.

The title compound was prepared by reaction of N-acetyl-tyrosinyl-valine and N-(N-allyloxycarbonyl-

25 4-benzyloxyprolinyl)-3-amino-4-oxobutanoic acid tertbutyl ester semicarbazone by reaction conditions reported for compound H, step A.

¹H NMR (500MH2, CD₃OD) δ 7.35-7.2 (m, 6H,, 7.0 (d, 2H), 6.65 (d, 2H), 4.85 (m, 1H), 4.6-4.45 (π, 4H), 30 4.3 (br. m, 1H), 4.15 (m, 1H), 3.7 (m, 1H), 2.95 (π,

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IH), 2.75-2.6 (m, 3H), 2.35 (m, 1H), 2.1 (m, 1H), 1.9 (s, 3H), 1.4 (s, 9H), 0.95 (d, 3H), 0.90 (s, 3H).

5

Step C. N-(N-Acetyl-tyrosinyl-valinyl-(4-benzyloxyprolinyl))-3-amino-4oxobutanoic acid.

benzyloxyprolinyl))-3-amino-4-oxobutanoic acid tertbutyl ester semicarbazone (270 mg) was dissolved into 10 ml of 25% trifluoroacetic acid in dichloromethane 10 and stirred at room temperature for 3 hours. The mixture was concentrated to give a solid residue. The residue was dissolved into a 10 ml mixture of methanol:acetic acid:37% formaldehyde (3:1:1) and stirred at room temperature for 1 hour. The mixture

N-(N-Acetyl-tyrosinyl-valinyl-(4-

15 was concentrated and the resulting residue purified by flash chromatography (SiO₂) eluting with dichloromethane/methanol/formic acid (100:5:0.5) to give 37 mg of the title compound.

 $^{1} H \ NMR \ (500 \ MHz, \ CD_{3}OD \ (existing as a 1:1)$ 20 mixture of diastereomers of the hemiacetal)) δ 7.4-7.25 $\{m, 5H\}, 7.0 \ (d, 2H), 6.65 \ (d, 2H), 4.65-4.05 \ (m, 7H), 3.75-3.4 \ (m, 2H), 3.05-2.3 \ (m, 5H), 2.2-1.95 \ (m, 2H), 1.90 \ (s, 3H), 1.0 \ (d, 3H), 0.95 \ (d, 3H).$

- (a) X = 0
- (b) $X = H_2$

(1S,9S) t-Butyl 6,10-dioxo-octahydro-9-(3-phenylpropionylamino)-6H-pyridazino[1,2-a]

- 5 [1,2]diazepine-1-carboxylate (44a). To a solution of (15,95) t-butyl 9-amino-6,10-dioxo-octahydro-6H-pyridazino [1,2-a][1,2]diazepine-1-carboxylate (690mg; 2.32mmol; GB 2128984) in dioxane (16ml) and water (4ml) at 0°C was added solid sodium bicarbonate (292mg;
- 10 3.48mmol) followed by dropwise addition of 3phenylpropionyl chloride (470mg; 2.78mmol). The mixture was stirred at room temperature for 2h then more sodium bicarbonate (200mg; 2.38mmol, and 3phenylpropionyl chloride (100mg; 0.6mmol) were added.
- 15 The mixture was stirred for a further 2h at room temperature, diluted with ethyl acetate (50ml), washed with saturated sodium bicarbonate (2 x 25ml) then dried (MgSO_d) and concentrated. The residue was purified by flash chromatography (0-50% ethyl acetate/chloroform)

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and finally crystallized by trituration with ether to afford 860mg (86%) of a white solid: mp. 137-138°C; $[\alpha]_D^{23}$ -95.1° (c 0.549, $\mathrm{CH}_2\mathrm{Cl}_2$); IR (KBr) 3327, 1736, 1677, 1664, 1536, 1422, 1156; $^1\mathrm{H}$ NMR (CDCl $_3$) δ 7.24 (5H, m), 6.50 (1H, d, J=7.5), 5.24 (1H, m), 4.90 (1H, m), 4.60 (1H, m), 3.44 (1H, m), 2.93 (2H, m), 2.84 (1H, m), 2.64 (1H, m), 2.54 (2H, m), 2.26 (2H, m), 1.70 (4H, m), 2.64 (2H, m), 2.26 (2H, m), 1.70 (4H, m), 2.64 (2H, m), 2.26 (2H, m), 1.70 (4H, m), 2.64 (2H, m), 2.26 (2H, m), 1.70 (4H, m), 2.64 (2H, m), 2.26 (2H, m), 1.70 (4H, m), 2.264 (2H, m), 2.264 (2H,

m), 1.70 (9H, s), MS(FAB, m/z): 430 (M+ + 1), 374.

10 (1s,9s) t-Butyl octahydro-10-oxo-9-(3-

242, 105, 91.

phenylpropionylamino)-6H-pyridazino-[1,2-a]
[1,2]diazepine-1-carboxylate (44b), was prepared from
(15,95) t-butyl 9-amino-octahydro-10-oxo-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxylate (Attwood

15 et al., <u>J. Chem. Soc. Perkin 1</u>, pp. 1011-19 (1986)) as for **44a**, to afford 810mg (81%) of a colorless oil: $\left[\alpha\right]_{\mathbf{D}}^{23}$ - 33.5° (c 0.545, CH₂Cl₂); IR (film) 3334, 2935, 1737, 1728, 1659, 1642; ¹H NNR (CDCl₃) δ 7.24 (5H, m), 6.75 (1H, d, \mathcal{J} =6.7), 5.27 (1H, m), 4.92 (1H, m), 3.39

20 (1H, m), 3.03 (4H, m), 2.55 (3H, m), 2.33 (1H, m), 2.17 (1H, m), 1.80 (5H, m), 1.47 (9H, s), 1.39 (1H, m).

MS(FAB, m/z): 416 (M⁺ + 1), 360, 211, 143, 97.

(1S,9S) 6,10-Dioxo-octahydro-9-(3-phenylpropionylamino)-6H-pyridazino[1,2-a]

25 [1,2]diazepine-1-carboxylic acid (45a). To a solution
 of (1S,9S) t-butyl 6,10-dioxo-octahydro-9-(3 phenylpropionylamino)-6H-pyridazino[1,2-a]
 (1,2]diazepine-1-carboxylate (44a) (800mg; 1.863mmol)
 in dry dichloromethane (5ml) at 0°C was added
30 trifluoroacetic acid (5ml). The solution was stirred
 at room temperature for 3h then concentrated. Dry

ether (10ml) was added to the residue then removed under vacuum. This process was repeated three times to afford a crystalline solid. The solid was triturated with ether and filtered to afford 590mg (85%) of a 5 white crystalline solid: mp. 196-197.5°C; [0]_D²³ -129.5° (c 0.2, CH₃0H); IR (KBr) 3237, 1729, 1688, 1666, 1633, 1574, 1432, 1285, 1205; ¹H NMR (CD₃0D) 5 8.28 (1H, d, J=7.4), 7.22 (5H, m), 5.32 (1H, dd, J=5.9, 2.9), 4.75 (1H, m), 4.51 (1H, m), 3.50 (1H, m), 3.01 (1H, m), 2.91 (2H, m), 2.55 (2H, m), 2.29 (3H, m), 1.95 (2H, m), 1.71 (2H, m). Anal. Calcd for Cl₁9H₂3N₃O₅: C, 61.12; H, 6.21; N, 11.25. Found: C, 60.80; H, 6.28; N, 10.97. MS(FAB, m/z) 374 (M⁴ + 1), 242, 105, 91.

(15,95) Octahydro-10-oxo-9-(3-phenylpropionylamino)-6Hpyridazino[1,2-a]-[1,2]diazepine-1-carboxylic acid
(45b), was prepared from (15,95) t-butyl octahydro-10oxo-9-(3-phenylpropionylamino)-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxylate (44b) by
the method described for compound 45a to afford 657mg
20 (96%) of 45b as a crystalline solid: mp. 198-202°C;
[α]_D²³ -86.2° (c 0.5, CH₃OH); IR (KBr) 3294, 2939, 1729,
1645, 1620, 1574, 1453, 1214; ¹H NMR (CD₃OD) δ 7.92
(1H, d, J=7.9), 7.20 (5H, m), 5.29 (1H, m), 4.90 (1H,
m), 3.47 (1H, m), 3.08 (2H, m), 2.90 (2H, m), 2.55 (3H,
25 m), 2.36 (1H, m), 1.81 (5H, m), 1.43 (2H, m), MS(FAB,

[3S,2R,S,(1S,9S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-octahydro-9-(3-phenylpropionylamino)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide (46a).

30 To a solution of (15,95) 6,10-dioxo-octahydro-9-13-phenyl-propionylamino)-6H-pyridazino[1,2-a]

m/z) 360 (M++1), 211,143,91,

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[1,2]diazepine-1-carboxylic acid (45a) (662mg; 1.773mmol) in dry dichloromethane (9ml) and dry dimethyl formamide (3ml) at room temperature was added bis(triphenylphosphine)palladium chloride (30mg) and 5 (3S, 2R, S) -3-allyloxycarbonylamino-2-benzyloxy-5oxotetrahydrofuran (Chapman, Bioorg, Med, Chem, Lett., 2, pp. 613-18 (1992)) (568mg; 1.95mmol) followed by dropwise addition of tri-n-butyltin hydride (1.19g; 4.09mmol). 1-Hydroxy-benzotriazole (479mg; 3.546mmol) 10 was added to the mixture and the mixture was cooled to 0°C before addition of 1-(3-dimethylaminopropyl)-3ethylcarbodiimide hydrochloride (408mg; 2.128mmol). The mixture was stirred at room temperature for 3.25h then diluted with ethyl acetate (50ml), washed twice 15 with dilute hydrochloric acid (20ml), twice with saturated sodium bicarbonate (20ml), once with brine then dried (MgSO4) and concentrated. The resulting oil was purified by flash chromatography (0-100% ethyl acetate/chloroform) to afford 810mg (81%) of 46a as a 20 mixture of anomers: mp. 92-94°C; IR (KBr) 3311, 1791, 1659, 1651, 1536; ¹H NMR(CDCl₃) 8 7.49, 6.56 (1H, 2d, J=6.7, 7.8), 7.29 (10H, m), 6.37, 6.18 (1H, 2d, J=7.7,7.6), 5.56, 5.34 (1H, d, s, J=5.2), 5.08-4.47 (6H), 3.18-2.80 (5H), 2.62-2.28 (5H), 2.04-1.53 (5H) 25 MS(FAB, m/z), 563 (M⁺ + 1), 328, 149, 91.

[3s,2R,s,(1s,9s)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-octahydro-10-oxo-9-(3-phenylpropionylamino)-6H-pyridazino[1,2-a]
[1,2]diazepine-1-carboxamide (46b), was prepared from
30 45b by the method described for 46a to yield 790mg
(96%) of a glass: m.p. 58-60°C; IR (KBE) 3316, 2940,
1793, 1678, 1641, 1523, 1453, 1120; H NMR (CDCl₃) 8

5 [3S(1S,9S)] 3-(6,10-Dioxo-octahydro-9-(3-

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7.28 (10H, m), 6.52, 6.42 (1H, 2d, J=7.2, 7.1), 5.53, 5.44 (1H, d, s, J=5.2), 5.35 (1H, m), 4.6-4.9, 4.34 (4H, m), 3.1-2.8 (6H, m), 2.6-2.1 (7H), 1.95-1.05 (5H). MS(FAB, m/z), 549 (M⁺ + 1), 400, 310, 279, 91.

phenylpropionylamino) -6H-pyridazino[1,2-a] [1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (47a). A mixture of [3S, 2R,S, (1S,9S)] N-(2-benzyloxy-5oxotetrahydrofuran-3-vl)-6,10-dioxo-octahydro-9-(3-10 phenylpropionylamino)-6H-pyridazino[1,2-a] [1,2]diazepine-1-carboxamide (46a) (205mg; 0.364mmol). 10% palladium on carbon (200mg) and methanol (20ml) was stirred under hydrogen at atmospheric pressure for 5h. The mixture was filtered then concentrated to yield 15 154mg (90%) of a glass: mp. 116-118°C; $[\alpha]_{\mathbf{p}}^{23}$ -140° (c 0.1, CH₃OH); IR (KBr) 3323 (br), 1783, 1731, 1658, 1539, 1455, 1425; ¹H NMR (CD₃OD) δ 7.21 (5H, m), 5.17 (1H, m), 4.73 (1H, m), 4.50 (2H, m), 4.23 (1H, m), 3.38 (1H, m), 3.06 (1H, m), 2.91 (2H, m), 2.73-2.18 (6H, m) 20 and 2.01-1.59 (5H, m). Anal. Calcd for $C_{23}H_{27}N_4O_7 + H_{2}O_7$: C, 56.32; H, 6.16; N, 11.42. Found: C, 56.29; H,

[3S(1S,9S)]3-(Octahydro-10-oxo-9-(3-

105, 91.

25 phenylpropionylamino) -6H-pyridazino-[1,2-a]
 [1,2]diazepine-1-carboxamido) -4-oxobutanoic acid (47b),
 was prepared from 46b by the method described for 47a.
 The residue was purified by flash chromatography (0-10
 methanol/chloroform) to afford 65mg (521) of a glass;
30 m.p. 87-90°C; [α]_D²³ -167.0° (c 0.1, methanol); IR
 (KBF) 3329, 2936, 1786, 1727, 1637; ¹H NMR (CD₂OD) 6

6.11; N, 11.25. MS(FAB, m/z) 473 (M+ 1), 176, 149.

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7.23 (5H, m), 5.29 (1H, m), 4.83 (1H, m), 4.59 (1H, d, J=3.6), 4.29 (1H, m), 3.3-3.0 (3H, m), 2.91 (2H, m), 2.70-2.34 (5H, m), 2.19 (2H, m), 1.75 (4H, m), 1.36 (2H, m). Anal. Calcd for $C_{23}H_{30}N_{4}O_{6}$ + 0.5H₂O: C, 59:09; 5 H, 6.68; N, 11.98. Found: C, 58.97; 6.68; N, 11.73. MS(FAB, m/z) 459 (M⁺ + 1), 310, 149, 105, 91.

t-Butyl N-2-(3-benzyloxycarbonylamino-1,2-dihydro-2-oxo-1- pyridyl)acetyl-3-amino-5-(2,6-dichloro-benzoyloxy)-4-oxo-pentanoate (56a). The acetic acid (55a) (WO 93 21213) in THF (2ml) was stirred at room temperature and treated with 1-hydroxybenzotriazole (60mg, 0.448mmol) and dimethylaminopropyl-3-ethylcarbodiimide hydrochloride (47mg, 0.246mmol). After 5 mins water (2 drops) was added and stirring continued for 20 minutes. Bis(triphenylphosphine) palladium TT chloride (6mg) was added followed by a solution of t-butyl 3-(allyloxycarbonylemino)-4-cxo-5-(2,6-dichlorobenzoyl-oxy)pentanoate (WO 93 16710) (103mg, 0.224mmol) in THF (lml). Tributyltin hydride

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- (0.09ml, 0.336mmol) was added dropwise over 1 hour at room temperature. The mixture was stirred for a further 3 hours and poured onto ethyl acetate, washed with 1M HCl, agueous NaHCO₂, brine, dried over MgSO₂
- 5 and concentrated *in vacuo*. The residue was triturated with pentane and the supernatant discarded. The remaining solid was purified by flash chromatography (50% ethyl acetate/hexame) to afford the title compound 92mg (63%) as a colorless oil: [cln²⁶ -29.6° (cl.1.)
- 10 CH_2Cl_2); IR (film) 3377, 3365, 3332, 3312, 1733, 1691, 1650, 1599, 1515, 1366, 1261, 1153, 1068, 747; 1_H NMR (CDCl₃) δ 8.09 (1H, d, J = 6.8), 7.84 (1H, s), 7.58 (1H, d, J = 8.3), 7.33 (8H, m), 7.02 (1H, dd, J = 6.9, 1.7), 6.33 (1H, t, J = 7.2), 5.20 (2H, s), 5.12 (2H,
- 15 m), 4.89 (1H, dt), 4.65 (2H, m), 2.80 (2H, m), 1.38 (9H, s).

t-Butyl N-2-(6-benzyl-1,2-dihydro-2-oxo-3-(3-phenylpropionyl)amino-1-pyridyl)acetyl-3-amino-5-(2,6-dichlorobenzyloxy)-4-oxo-pentanoate (56b), was prepared

- 20 by the method described for (56a) which afforded the title compound (66%) as a colorless oil: IR (film) 3364, 3313, 1738, 1688, 1648, 1600, 1566, 1514, 1433, 1369, 1254, 1152; ¹H NMR (CDCl₃) δ 8.40 (1H, d, J 7.6), 8.30 (1H, s), 7.28 (13H, m), 6.20 (1H, d, J = 7.6),
- 25 5.12 (2H, q), 4.86 (1H, m), 4.65 (2H, q), 4.06 (2H, s), 3.07--2.61 (6H, m), 1.39 (9H, s).

N-2 (3-Benzyloxycarbonylamino-1,2-dihydro-2-oxo-1pyridyl)acetyl-3-amino-5-(2,6-dichlorobenzovloxy)-4oxo-pentanoic acid (57a; Q). The ester 56a (210mg, 0.356mmol) in dichloromethane (0.5ml) was cooled to 0°C 5 and treated with trifluoroacetic acid (0.5ml), stirred and warmed to 20°C over 30 minutes. The solution was evaporated to dryness under reduced pressure, redissolved in dichloromethane and concentrated (x3). The residue was triturated with ethyl acetate and 10 diluted with ether to afford the title compound 162mg (85%) as a colorless solid: m.p. 165-8°C (decomposition); $[\alpha]_{\mathbf{D}}^{23}$ -38.8° (c 0.1, CH₃OH); IR (KBr) 3332, 3275, 1723, 1658, 1649, 1597, 1581, 1562, 1526, 1432, 1385, 1258, 1218, 1206; ¹H NMR (dz-DMSO) & 3.96 15 (1H, d, J = 7.3), 8.34 (1H, s), 7.85 (1H, dd, J = 7.3). 7.58 (3H, m), 7.35 (5H, m), 6.29 (1H, t, J = 7.3), 5.26 (2H, m), 5.15 (2H, s), 4.69 (3H, m), 2.75 (2H, m). Anal. Calcd. C27H23N3OgClo: C, 53.66; H, 3.84; N, 6.95. Found: C, 53.36; H, 3.90; N, 6.81, M.S. (+ FAB:: 604 20 (M⁺ + 1), 285, 241, 195, 173, 149, 91.

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N-2-(6-Benzyl-1,2-dihydro-2-oxo-3-(3-phenylproplonyl)
amino-1-pyridyl)acetyl-3-amino-5-(2,6-dichlorobenzoyloxy)-4-oxo-pentanoic acid (57b; P), was prepared
by the method described for 57a which afforded the
5 title compound (78%) as colorless crystals: m.p. 116120°C (decomposition); [\alpha]_0^{26} -41.1° (c 0.1, CH3OH); IR
(KBr) 3299, 1739, 1715, 1689, 1666, 1645, 1598, 1563,
1518, 1432, 1209, 1151; \frac{1}{4} H NMR (d_6-DMSO) & 9.24 (1H,
s), 8.88 (1H, d, J = 7.6), 8.18 (1H, d, J = 7.7), 7.60

10 (3H, m), 7.26 (10H, m), 6.06 (1H, d, J = 7.7), 5.23
(2H, ABg), 4.69 (3H, m), 3.93 (2H, s), 2.78 (6H, m).

Anal. Calcd. for $C_{35}H_{31}N_{30}gCl_2$. H_2O : C, 59.16; H, 4.68; N, 5.91. Found: C, 59.38; H, 4.53; N, 5.84. M.S. (+ FAB); 694, (Cl=35, 37), (M⁺ + 1); 692 (Cl=35, 35), (M⁺ 15 + 1).

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(a)
$$R^1 = OCH_3$$
, $R^2 = H$
(b) $R^1 = H$, $R^2 = OCH_3$

(b)
$$R^1 = H$$
, $R^2 = OCH_3$

7-Methoxybenzoxazole (65a). A mixture of 2-nitro-6methoxyphenol (2.62g, 15.5mmol) (EP 333176) and 10% 5 Palladium on carbon (130mg) in ethanol (50.0ml) was stirred under an atmosphere of H2 for 75min. The

mixture was filtered through Celite® then immediately treated with p-toluenesulphonic acid (32.0mg) and triethylorthoformate (6.45ml, 38.8mmol) then heated under reflux under an atmosphere of N_2 . After 20h p-

- 5 toluenesulphonic acid (30.0mg) and triethylorthoformate (6.45ml, 38.8mmol) were added. After a total of 44h heating, the reaction was allowed to cool and reduced in vacuo. The resulting residue was purified by flash chromatography (25:75 ethyl acetate/hexane) to give
- 10 1.97g (85%) of the title compound as a yellow solid: m.p. $28-31^{\circ}\text{C}$; IR (film) 1629, 1497, 1434, 1285, 1097; ^{1}H NMR (CDCl₃) δ 8.09 (1H, s), 7.40 (1H, d, J = 8.0), 7.28 (1H, t, J = 8.0), 6.89 (1H, d, J = 8.0), 4.02 (3H, s); ^{13}C NMR (CDCl₃) δ 152.84, 145.82, 142.50, 139.99,
- 15 125.75, 113.42, 108.80, 56.97. Anal. Calcd. for
 C₈H₇N₂O₂. 0.1H₂O: C, 63.65; H, 4.81; N, 9.29. Found:
 C, 63.43, H, 4.88, N, 9.05. M.S. (+ FAB); 150 (M⁺ + 1).

- (25:75 ethyl acetate/hexame) to give 2.0g (91°) of the title compound as a white crystalline solid: m.p. 72-74°C; IR (KBr) 3089, 1619, 1610, 1503, 1496, 1322, 1275, 1090, 1071, 780, 741; H NMR (CDCl₃) & 8.02 (18,
- 30 s), 7.32 (1H, t, J=8.0), 7.18 (1H, d, J=8.0), 6.81 (1H, d, J=8.0), 4.04 (3H, s). Anal. Calcd. for

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 $C_8H_7NO_2$: C, 64.42; H, 4.73; N, 9.39. Found: C, 64.40; H, 4.84; N, 9.31; m/z (EI) 149 (M⁺ + 1, 100%).

(3S, 4R,S) t-Butyl N-(allyloxycarbonyl)-3-amino-4-hydroxy-4-(2-(7-methoxybenzoxazolyl))butanoate (66a).

- 5 To a stirred solution of 7-methoxybenzoxazole 65a (540.6mg, 3.68mmol) in anhydrous THF (18.5ml) at -78°C under N_2 was added 1.56M n-butyl lithium in hexanes (2.47ml, 3.86mmol) dropwise, to produce a yellow colored solution. After stirring at -78°C for 20 min,
- 10 dry MgBr $_2$ OEt $_2$ (1.045g, 4.05mmol) was added as a solid. The resulting heterogeneous mixture was warmed to $-45\,^{\circ}$ C and stirred for 15min. The reaction mixture was then recooled to $-78\,^{\circ}$ C and a solution of (S)-Alloc-Asp(t-Bu)H (946.4mg, 3.68mmol) in THF (18.5ml) was added
- 15 dropwise. The reaction was stirred at -78°C for 30min, warmed to 0°C and stirred for lh. The resulting homogeneous reaction was warmed to room temperature and stirred for 16h. The reaction was quenched with 5% sodium bicarbonate (3.5ml) then THF was removed in
- 20 vacuo. The resulting aqueous residue was extracted with methylene chloride (x6). The combined extracts were washed with brine, dried (MgSO₄), filtered and reduced in vacuo to give 1.8g of crude product. Flash chromatography (40:60 ethyl acetate/hexane) gave 1.21g
- 30 (2H, m⁺, 1.45 (9H, s), 1.41 (9H, s); ¹³C NMR (CDCl₃) 8 171.18, 171.09, 165.80, 165.30, 156.71, 156.60, 145.65, 142.76, 142.71, 140.82, 140.72, 133.23, 125.81, 125.72,

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118.41, 118.21, 113.07, 112.87, 108.95, 82.16, 70.28, 69.98, 66.52, 66.39, 57.03, 52.57, 52.29, 37.83, 36.86, 28.65. Anal. Calcd. for C₂₀H₂₆N₂O₇. 0.6H₂O: C, 57.57; H, 6.57; N, 6.72. Found: C, 57.49, H, 6.34, N, 6.60. 5 M.S. (+ FAB); 407 (M⁺ + 1); 351, 307, 154.

- (3S, 4R,S) t-Butyl N-(allyloxycarbonyl)-3-amino-4hydroxy-4-(2-(4-methoxybenzoxazolyl))butanoate (66b), was prepared according to the method described for 66a which afforded 1.29g (26%, 68% based on recovered
- 10 starting material) of the title compound as an oil and as a mixture of diastereoisomers at C-4: IR (CH₂Cl₂) 3400, 1725, 1625, 1505, 1369, 1354, 1281, 1263, 1226, 1158, 1092, 1048; 1 H NMR (CDCl₃) δ 7.34-7.24 (1H, m), 7.16 (1H, d, J = 8.2), 6.79 (1H, d, J = 7.9), 6.00-5.50
- 15 (2H, m), 5.30-5.05 (3H, m), 4.70-4.35 (4H, m), 4.02 (3H, s), 2.90-2.45 (2H, m), 1.45-1.41 (9H, 2 x s). Anal. Calcd. for $C_{20}H_{26}N_{2}O_{7}$. 0.4 $H_{2}O$: C, 58.07; H, 6.53; N, 6.77. Found: C, 58.09; H, 6.41; N, 6.63. M.S. (+ FAB); 407 (M^{+} + 1, 88%); 351 (100).
- 20 (3S, 4R,S) t-Butyl N-(N-acetyl-(S)-(O-tert-butyl-tyrosinyl)-(S)-valinyl-(S)-alaninyl)-3-amino-4-hydroxy-4-(2-(7-methoxybenzoxazolyl))butanoate (67a). To a stirred solution of the benzoxazole 66a (481.9mg, 1.19mmol) and Ac-Tyr(tBu)-Val-Ala-OH (586.3mg,
- 25 1.30mmol) in methylene chloride (3.5ml) and DMF (3.5ml) was added bis(triphenylphosphine) palladium (II) chloride (18.0mg), followed by tributyltinhydride (0.80ml, 2.96mmol) dropwise. Hydroxybenzotriazole (320.4mg, 2.37mmol) was added and the mixture cooled to
- 30 0°C. 1-Ethyl-3-[3-(dimethylamino)propyl]carbodizmide hydrochloride (278.2mg, 1.42mmol) was added and the

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mixture was allowed to warm to room temperature and stirred for 16.5h. The reaction was diluted with ethyl acetate and washed twice with 1M sodium hydrogensulphate, twice with saturated sodium

- 5 bicarbonate, water, and brine. The organic layer was dried (MgSO4), filtered and reduced in vacuo to yield 2.0g of crude product. Flash chromatography (95:5 methylene chloride/methanol) gave 844.0mg (94%) of the title compound as a white solid: m.p. 205°C; IR (KBr)
- 10 3399, 3304, 2977, 1729, 1643, 1506, 1367, 1290, 1161; ¹H NMR (d₆-DMSO) δ 8.24-7.78 (4H, m), 7.43-7.32 (2H, m), 7.23 (2H, d, J = 8.5), 7.16-7.07 (1H, m), 6.93 (2H, d, J = 8.5), 6.52, 6.40 (1H, 2 x d, J = 5.5, J = 5.0), 5.03, 4.78-4.49, 4.45-4.16 (5H, brt, 2 x m), 4.05, 4.04
- 15 (3H, 2 x s), 3.08-2.35 (14H, m), 2.11-1.89 (1H, m), 1.83 (3H, s), 1.49-1.32, 1.15, 1.0-0.81 (27H, s, 2 x m, J = 7.0); ¹³C NMR (d₆-DMSO) δ 175.55, 175.18, 173.88, 173.75, 173.05, 169.23, 157.28, 148.55, 146.16, 143.21, 136.63, 133.55, 128.87, 127.17, 115.78, 111.92, 84.02,
- 20 81.50, 71.40, 61.15, 60.05, 57.79, 53.39, 51.62, 43.76, 40.52, 34.58, 32.52, 31.60, 26.35, 23.11, 22.71, 21.76. Anal. Calcd. for C39H55N5O10. 0.5H2O: C, 61.40; H, 7.40; N, 9.18. Found: C, 61.43; H, 7.31; N, 9.07. M.S. (+ FAB); 754 (M+ + 1); 698, 338, 267.
- 25 (3S, 4R,S) t-Butyl N-(N-acetyl-(S)-(O-tert-butyltyrosinv1) - (S) -valinyl-(S) -alaninyl) -3-amino-4-hydroxy-4-(2-(4-methoxybenzoxazolyl))butanoate (67b), was prepared according to the method described for 67a which afforded 1.05g (94%) of the title compound as a 30 fine white powder: m.p. 210-213°C (dec); IR (KBr) 3284, 2977, 1736, 1691, 1632, 1536, 1505, 1452, 1392, 1367, 1258, 1236, 1161, 1091; ¹H NMR (d₆-DMSC) δ 8.20-

7.75 (4H, m), 7.40-7.10 (4H, m), 7.00-6.80 (3H, m), 6.45, 6.34 (1H, 2 x d, J = 5.3, J = 5.0), 5.00-4.10 (5H, m), 4.00, 3.99 (3H, 2 x s), 3.00-2.25 (4H, m), 1.95 (1H, m), 1.78 (3H, s), 1.39-0.80 (27H, m). And 1. 5 Calcd. for $C_{33}H_{55}N_{5}O_{10}$. 0.5H₂O: C, 61.40; H, 7.40; N, 9.18. Found: C, 61.58; H, 7.38; N, 8.91. M.S. (+ FAB); 754 (M⁺ + 1, 30%); 72 (100).

(3S) t-Butyl N-(N-acetyl-(S)-(O-tert-butyl-tyrosinyl)-(S)-valinyl-(S)-alaninyl)-3-amino-4-(2-(7-

- 10 methoxybenzoxazolyl))-4-oxobutanoate (68a). The DessMartin reagent (1.082g, 2.55mmol) (Ireland et al., J.
 Org. Chem., 58, p. 2899 (1993); Dess et al., J. Org.
 Chem., 48, pp. 4155-4156 (1983)) was added to a stirred
 suspension of the alcohol 67a (641.0mg, 0.85mmol) in
- 15 methylene chloride (46.0ml). The resulting mixture was stirred for lh before being partitioned between saturated sodium thiosulphate: saturated sodium bicarbonate (1:1, 86.0ml) and ethyl acetate (86.0ml). The resultant organic phase was washed in turn with
- 20 saturated sodium thiosulphate: saturated sodium bicarbonate (1:1), saturated sodium bicarbonate, and brine. The organic phase was dried (MgSO₄), filtered and reduced in vacuo to give 660.0mg of crude product. Flash chromatography (94:6 methylene chloride/methanol)
- 25 gave 636.0mg (100%) of the title compound as a wnite solid: m.p. 209°C; [α]_D²⁴ -21.8° (c 0.16, methanol); IR (KBr) 3395, 3294, 2977, 1722, 1641, 1535, 1505, 1161; ¹H NMR (CDCl₃) 8 8.43-8.16 (1H, m), 7.97-7.62 (2H, m), 7.49-7.14 (3H, m), 7.08-6.95 (3H, m), 6.89-6.73 (2H.
- 30 m), 5.81-5.68 (1H, m), 5.16-4.86 (2H, m), 4.53 (1H, brt), 4.03 (3H, s), 3.16-2.84 (4H, m), 2.1i-1.84 (4H, m), 1.46-1.14 (21H, m), 0.92-0.78 (6H, m); ¹³C NMR

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(CDC1₃) & 186.28, 173.39, 171.90, 171.19, 171.03, 169.89, 156.43, 154.75, 146.32, 142.88, 140.98, 132.31, 130.54, 126.98, 124.73, 114.95, 111.42, 82.44, 78.71, 58.92, 57.20, 54.91, 53.47, 48.77, 39.43, 38.15, 32.79, 54.42, 28.60, 23.55, 20.27, 19.70, 19.34. M.S. (+ FAB); 752 (M + 1); 696, 336, 265.

(3S) t-Butyl N-(N-acetyl-(S)-(O)-tert-butyl-tyrosinyl)-(S)-valinyl-(S)-alaninyl)-3-amino-4-(2-(4-methoxybenzoxazolyl))-4-oxobutanoate (68b), was

10 prepared according to the method described for the ketone 68a which afforded 420mg (55%) of the title compound as a white solid: m.p. 211-213°C (dec); (α)_D2⁴ -23.9° (c 0.82, methanol); IR (KBr) 3277, 3075, 1723, 1690, 1632, 1530, 1506, 1392, 1366, 1269, 1234, 1160, 1094; ¹H NMR (CDCl₃) δ 8.15 (1H, brs), 7.7 (2H, brs), 7.46 (1H, t, J = 8.3), 7.24 (2H, d, J = 8.3), 7.10 (1H, brs), 7.03 (2H, d, J = 8.3), 6.83 (3H, m), 5.74 (1H, q, J = 6.9), 5.00 (2H, m), 4.51 (1H, t, J = 7.0), 4.07 (3H, s), 3.20-2.95 (4H, m), 2.00 (4H, m), 1.42 (3H, d, 2 = 6.8), 1.35 (9H, s), 1.23 (9H, s), 0.86 (6H, d, J = 6.7), M.S. (+ FAB); 752 (M* + 1, 7%); 72 (100).

(3S) N-(N-Acetyl-(S)-tyrosinyl-(S)-valinyl-(S)alaninyl)-3-amino-4-(2-(7-methoxybenzoxazolyl))-4oxobutanoate (69a; R). A solution of the ester 68a
25 (600.0mg, 0.80mmcl) in a 1:1 mixture of methylene
chloride and trifluoroacetic acid (65.0ml) was stirred
for 1h under a dry atmosphere of N₂. The solution was
then reduced in vacuo, taken up in ether and reduced
again. This process was repeated six times to afford
30 the crude product as an off white solid. Flash
chromatography (gradient 95:5 to 80:20 methylene

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chloride/methanol) gave 420.8mg (83%) of the title compound as a hygroscopic white solid. The product existed as a mixture of three isomers in CD₃OD, consisting of the keto form (c 50%), and its acycloxy keto form (two isomers at C-4, c 50%): m.p. decomposes above 150°C; [α]₀²⁴-33.2° (c 0.17, methanol); IR (KBr) 3300, 1715, 1658, 1650, 1531, 1517, 1204;

14 NMR (CD₃OD) δ 7.46-7.19 (2H, m), 7.16-6.91 (3H, m), 6.70-6.59 (2H, m), 5.62-5.49 (1H, m), 5.00-4.72 (1H, 0 obscurred m), 4.69-4.51 (1H, m), 4.49-4.08 (2H, m), 4.05-3.89 (3H, m), 3.16-2.47 (4H, m), 2.05-1.78 (4H, m), 1.41-1.11, 1.05-0.70 (9H, 2 x m). Anal. Calcd. for C₃₁H₃₇N₅O₁₀. 3H₂O: C, 53.67; H, 6.25; N, 10.10. Found: C, 53.76; H, 5.56; N, 10.28. M.S. (+ FAB); 640 (M + 1); 435, 147.

(3S) t-Butyl N-(N-acetyl-(S)-tyrosinyl-(S)-valinyl-(S)alaninyl) -3-amino-4-(2-(4-methoxybenzoxazolyl))-4oxobutanoate (69b; S), was prepared according to the method described for the acid 69a which afforded the 20 hygroscopic title compound 252mg (96%). The product existed as a mixture of three isomers in CD2OD, consisting of the keto form, and its acycloxy ketal form (two isomers at C-4). The product existed as a single isomer in d-6 DMSO: m.p. 200-203°C (dec.); 25 (a)_n²⁴ -38.0° (c 0.23, methanol); IR (KBr) 3289, 2968, 1718, 1713, 1658, 1634, 1548, 1517, 1506, 1461, 1453. 1393, 1369, 1268, 1228, 1174, 1092; ¹H NMR (de-DMSO) & 9.20 (1H, brs), 8.71 (1H, d, J = 6.2), 8.10 (2H, m), 7.83 (1H, d, J = 8.7), 7.61 (1H, t, J = 8.2), 7.46 (1H, 30 d, J = 8.2), 7.08 (3H, m), 6.65 (2H, d, J = 8.3), 5.50 (IH, q, J = 6.5), 4.50 (IH, m), 4.37 (IH, m), 4.20 (IH, m , 4.05 (3H, s), 3.09-2.77 (4H, m), 1.94 (1H, m), 1.79

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(3H, s), 1.23 (3H, d, J = 7.0), 0.82 (6H, m). Anal. Calcd. for $C_{31}H_{37}N_5O_{10}$. 1.5 H_2O : C, 55.85; H, 6.05; N, 10.51. Found: C, 55.21; H, 5.69; N, 10.13. M.S. (+ FAB); 640 (M^4 + 1, 22%); 107 (100).

5 3(S)-(Allyloxycarbonyl)-amino-4-[(2,6-dichlorophenyl) -oxazol-2-vl]-4(R,S)-hydroxy-butyric acid tertbutyl ester (99). A solution of 5-(2,6-Dichlorophenyl) oxazole (2.71g, 12.7mmol; prepared by a similar method described in Tet. Lett. 23, p. 2369 10 (1972)) in tetrahydrofuran (65mL) was cooled to -78 °C under a nitrogen atmosphere. To this solution was added n-butyl lithium (1.5M solution in hexanes, 8.5mL, 13.3mmol) and stirred at -78 °C for 30min. Magnesium bromide etherate (3.6g, 13.9mmol) was added and the 15 solution was allowed to warm to -45 °C for 15min. The reaction was cooled to -78 °C and aldehyde 58 (3.26g, 12.7mmol; Graybill et al., Int. J. Protein Res., 44, pp. 173-182 (1993)) in tetrahydrofuran (65mL) was added dropwise. The reaction was stirred for 25min., then 20 allowed to warm to -40 °C and stirred for 3h, and then at room temperature for 1h. The reaction was quenched with 5% NaHCO3 (12mL) and stirred for 3h. The tetrahydrofuran was removed in vacuo and the resulting residue was extracted with dichloromethane. The 25 organic layer was washed with saturated sodium chloride solution and dried over magnesium sulfate, filtered,

and concentrated to yield 6.14g of the title compound.

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Purification gave 4.79g (80%) of 99: ¹H NMR (CDCl₃) δ 1.45(s, 9H), 2.7-2.5(m, 2H), 2.8(dd, 1H), 4.2, 4.4(2 x d, 1H), 4.7-4.5(m, 3H), 5.35-5.1(m, 2H), 5.6, 5.7(2 x d, 1H), 6.0-5.8(m, 1H), 7.2(d, 1H), 7.3(m, 1H), 7.4(m, 5 2H).

a R = H

b R = COCH2CH2Ph

 $c R = CH_2Ph$

[2-0xo-3(S)-(3-phenylpropionylamino)-2,3,4,5
10 tetrahydro-benzo[b][1,4]diazepin-1-yl]acetic acid
methyl ester (104a). Anhydrous hydrogen chloride was
bubbled into a solution of (3(S)-tertbutoxycarbonylamino-2-oxo-2,3,4,5-tetrahydro-benzo[b]
[1,4]diazepin-1-yl]acetic acid methyl ester (103, 1g,

15 2.96 mmol) in 25 ml of ethyl acetate for 2 minutes then stirred for 1 hour at room temperature. The reaction was evaporated to give 2-oxo-3(S)-amino-2,3,4,5-

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tetrahydrobenzo[b][1,4]diazepin-1-yl acetic acid methyl ester hydrochloride as a white solid. The hydrochloride salt and hydrocinnamic acid (0.47 c. 3.15 mmol) were dissolved into 20 ml of 5 dimethylformamide and cooled to 0 °C. Diisopropylethylamine (1 ml, 5.72 mmol) was added to the solution followed by the addition of Nhydroxybenzotriazole and 1-(3-dimethylaminopropyl)-3ethylcarbodiimide hydrochloride. After stirring for 18 10 hours at room temperature, the mixture was diluted with 150 ml of ethyl acetate and washed with 10% sodium hydrogen sulfate, 10% sodium bicarbonate, and brine. The organic layer was dried over anhydrous sodium sulfate, filtered, and evaporated to a crude solid that 15 was purified by flash chromatography eluting with 7:3 ethyl acetate/dichloromethane to afford 600 mg (55%) of the title compound as a white solid. 1 H NMR (CDCl₂) δ 7.3-6.85 (9H,m), 6.55-6.0 (1H, d), 4.88-4.82 (1H, m), 4.72-4.65 (1H, d), 4.28-4.22 (1H, m), 3.95-3.9 (1H, m),

(3(S)-(3-Phenylpropionylamino)-2-oxo-2,3,4,5-tetrahydrobenzo[b][1,4]diazepin-1-yl)acetic acid (105a). (3(S)-(3-Phenylpropionylamino)-2-oxo-2,3,4,5-

20 3.78 (3H, s), 3.65 (1H, br. s), 3.28-3.2 (1H, m), 2.95-

2.84 (2H, m), 2.55-2.4 (2H, m),

25 tetrahydro-benzo(b) [1,4]diazepin-1-yllacetic acid
methyl ester (104a) was dissolved in 90° methanol.
Lithium hydroxide hydrate was added to the reaction and
the reaction was stirred at room temperature for 4 h.
The reaction was evaporated in vacuo to give a white
30 solid. This was dissolved in 20 ml of water and
acidified to pH 5 and extracted with ethyl acetate to
afford 304 mg (88%) of the title compound as a white

solid. 1 H NMR (CDCl₃) δ 7.5-6.9 (11H, m), 4.92-4.8 (1H, m), 4.7-4.58 (1H, d), 4.38-4.25 (1H, d), 3.88-3.78 (1H, m), 3.45-3.25 (1H, m), 3.05-2.85 (2H, m), 2.55-2.45 (2H, m).

- 5 4-0xo-3(S)-(2-[2-oxo-3(S)-(3-phenylpropionylamino)-2,3,4,5-tetrahydro-benzo[b][1,4]diazepin-1ylacetylamino)butyric acid (106a). N-[1-(2-Benzyloxy-5-oxotetrahydrofuran-3-vlcarbamovl-methvl)-2-oxo-2,3,4,5-tetrahydro-1H-benzo[b][1,4]dlazepin-3-yl]-3-10 phenylpropionamide was prepared from 105a by the procedure used to prepare compound H (stepA) to afford 390 mg (93%) of the product as diastereomers. ¹H NMR (CD₃OD) δ 7.58-7.22 (14H, m), 5.78-5.73 (0.5 H, d), 5.64 (0.5 H, s), 5.0-4.72 (4H, m), 4.54-4.42 (2H, m), 3.82-15 3.76 (0.5 H, m), 3.68-3.62 (o.5 H, m), 3.28-3.21 (0.5H, m), 3.19-3.12 (0.5H, m), 3.07-2.98 (2H, m), 2.76-2.48 (4H, m). The resulting product was converted to 106a by the method described to prepare compound H (StepD) to 20 afford the title compound as a white solid (17%): 1H NMR (CD₃OD) δ 7.54-6.98 (9H, m), 5.58-5.44 (1H, m), 4.8-
 - [2-0xo-5-(3-phenylpropionyl)-3(S)-(3-

2.25 (5H, m).

25 phenylpropionylamino)-2,3,4,5tetrahydrobenzo[b][1,4]diazepin-1-yl]acetic acid methyl
 ester (104b). Anhydrous hydrogen chloride was bubbled
 into a solution of (3(S)-tert-butoxycarbonylamino-2 oxo-2,3,4,5-tetrahydro-benzo[b][1,4]diazepin-130 yl]acetic acid methyl ester (103, 19, 2.86mmol) in 25

4.2 (4H, m), 3.96-3.3 (2H, m), 3.30-3.05 (1H, m), 2.98-

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ml of ethyl acetate for 2 minutes then stirred for 1 hour at room temperature. The reaction was evaporated to give 2-oxo-3(S)-amino-2,3,4,5-tetrahydrobenzo[b][1,4]diazepin-1-yl acetic acid methyl

testranydrobenzoloj[[1,4]d_azephn=1-yl acetic acid methy
sester hydrochloride as a white solid.

The hydrochloride salt was suspended into 20 ml of dichloromethane and cooled to 0 °C. Triethylamine (1.6 ml, 11.5 mmol) was added to the suspension followed by the dropwise addition of dihydrocinnamoyl chloride (0.9

10 ml, 6 mmol). The mixture was warmed to room temperature and stirred for 18 hours. The mixture was diluted with 25 ml of dichloromethane and washed twice with 50 ml of water and once with 50 ml of brine. The organic layer was dried over anhydrous sodium sulfate,

15 filtered, and evaporated to give a viscous, yellow oil that was purified by flash chromatography eluting with 1:1 ethyl acetate/dichloromethane to afford 1.35 g (92%) of the title product as a white solid. $^{1}\mathrm{H}$ NMR (CDCl₃) δ 7.45-7.02 (14 H, m), 6.37-6.32 (1H, d), 4.78-

20 4.72 (1H, m), 4.52-4.3 (3H, m), 3.82-3.77 (1H, m), 3.74 (3H, s), 3.03-2.87 (4H, m), 2.58-2.45 (2H, m), 2.45-2.35 (1H, m), 2.25-2.16 (1H, m).

[2-0xo-5-(3-phenylpropionyl)-3-(3(S)-phenylpropionylamino)-2,3,4,5-

25 tetrahydrobenzo[b][1,4]diazepin-1-yl]acetic acid
(105b). [2-0xo-5-(3-phenylpropionyl)-3-(3-phenylpropionylamino)-2,3,4,5tetrahydrobenzo[b][1,4]diazepin-1-yl]acetic acid methyl
ester (104b; 680 mg, 1.32 mmol) was hydrolyzed by the
30 procedure used to hydrolyze 105a to afford 645 mg (98.)
of the title compound as a white solid. ¹H NMR 'CDCl₃)
& 7.56 (HH, br. s), 7.5-7.42 (HH, m), 7.35-6.95 '14H.

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m), 4.95-4.88 (1H, m), 4.64-4.55 (1H, d), 4.54-4.45 (1H, t), 4.15-4.05 (1H, d), 3.75 (1H, m), 3.05-2.75 (4H, m), 2.58-2.45 (2H, m), 2.45-2.28 (1H, m), 2.25-2.14 (1H, m).

- 5 2-Oxo-3(S)-(2-[2-oxo-5-(3-phenylpropionyl)-3(S)-(3-phenyl-propionyl-amino)-2,3,4,5tetrahydrobenzo[b][1,4]diazepin-1-yl]
 acetylamino}butyric acid (106b). [2-Oxo-5-(3-phenylpropionyl)-3-(3-phenylpropionylamino)-2,3,4,5-
- 10 tetrahydrobenzo[b][1,4]diazepin-1-yl]acetic acid and 3-amino-4-oxobutyric acid tert-butylester semicarbazone were coupled by the procedure used in the preparation of compound K (step A) to give 350 mg (85%) of a white solid. $^{1}{\rm H}$ NMR (CDCl3) δ 9.05 (1H, br. s), 7.58-7.55
- 15 (1H,d), 7.5-7.35 (1H, m), 7.35-6.95 (14 H, m), 6.75-6.72 (1H, d), 6.25 (1H, br. s), 5.25 (1H, br. s), 4.95-4.88 (1H, m), 4.8-4.72 (1H, m), 4.55-4.4 (2H, m), 3.92-3.88 (1H, d), 3.73-3.68 (1H, m), 2.95-2.8 (4H, m), 2.8-2.72 (1H, m), 2.62-2.55 (1H, m), 2.55-2.45 (2H, m),
- 20 2.4-2.32 (1H, m), 2.2-2.12 (1H, m), 1.45 (9H, s).
 4-0xo-3-{2-[2-oxo-5-(3-phenylpropionyl)-3-(3-phenyl-propionyl -amino)-2,3,4,5tetrahydrobenzo[b][1,4]diazepin-1-yl]-acetylamino]butvrc acid tett-butvl ester semicarhazone was
- 25 deprotected as described in the preparation of compound K (step C) to give 118 mg $(47\,\text{m})$ of the title compound as a white solid. ^{1}H NMR (CD₃OD) δ 7.48-6.95 (14 H, m), 4.65-4.15 (6H, m), 3.5-3.4 (1H, m), 2.85-2.72 (4H, m), 2.65-2.5 (1H, m), 2.5-2.34 (3H, m), 2.34-2.15 (2H, m).
- 30 [5-Benzyl-2-oxo-3(S)-(3-phenylpropionylamino)-2,3,4,5-tetrahydro-benzo[b][1,4]diazepin-1-yl]acetic acid

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methyl ester (104c). [2-0xo-3-(3phenylpropionylamino) -2,3,4,5-tetrahydrobenzo-[b][1,4]diazepin-1-vl]acetic acid methyl ester (104a; 500 mg, 1.31 mmol), calcium carbonate (155 mg, 1.58 5 mmol), and benzyl bromide (170 ul, 1.44 mmol) were taken into 10 ml of dimethylformamide and heated to 80 °C for 8 hours. The mixture was diluted with 150 ml of ethyl acetate and washed 4 times with 50 ml of water. The organic layer was dried over anhydrous sodium 10 sulfate, filtered, and evaporated to give a viscous, yellow oil that was purified by flash chromatography eluting with dichloromethane/ethvl acetate (8:2) to give 460 mg (75%) of the title compound as a white solid. ¹H NMR (CDCl₃) δ 7.34-7.05 (14 H, m), 6.32-6.28 15 (1H, d), 4.84-4.76 (1H, d), 4.76-4.70 (1H, m), 4.43-4.37 (1H, d), 4.26-4.18 (1H, d), 4.06-4.00 (1H, d), 3.79 (3H, s), 3.45-3.37 (1H, m), 3.02-2.95 (1H, m), 2.90-2.82 (2H, m), 2.5-2.34 (2H, m).

[5-Benzyl-2-oxo-3(S)-(3-phenylpropionylamino)-2,3,4,5-20 tetrahydro-benzo[b][1,4]diazepin-1-yl]acetic acid (105c) was prepared by the hydrolysis of the ester (102c) by the procedure reported in Example 105a to give 450 mg (98%) of the title compound as a white solid: ¹H NMR (CD₃OD) δ 7.5-7.05 (14 H, m), 6.4 (1H, 25 br. s), 4.85-4.55 (2H,m), 4.5-4.21 (2H, m), 4.12-3.92 (1H, d), 3.45-3.3 (1H, m), 3.1-2.8 (3H, m), 2.55-2.28 (3H, m).

3(s)-{2-[5-Benzyl-2-oxo-3-(3(s)-phenylpropionylamino)-2,3,4,5-tetrahydrobenzo[b][1,4]diazepin-1-yl]30 acetylamino}-4-oxobutyric acid (106c). [5-Benzyl-2-

oxo-3(S)-(3-phenylpropionylamino)-2,3,4,5-tetrahydrobenzo[b[1,4]diazepin-1-yl]acetic acid and 3(S)-amino-4-oxobutyric acid tert-butylester semicarbazone were coupled by the procedure used in the preparation of

- 5 compound K (step A) and to afford 260 mg (85%) of a white solid: ¹H NMR (CD₃OD) δ 7.35-7.0 (15 H, m), 4.94-4.88 (1H, m), 4.68-4.58 (1H, d), 4.57-4.52 (1H, m), 4.41-4.34 (1H, d), 4.3-4.23 (1H, d), 4.1-4.04 (1H, d), 3.18-3.11 (1H, m), 3.09-2.98 (1H, m), 2.78-2.72 (2H,
- 10 t), 2.65-2.57 (1H, m), 2.42-2.33 (3H, m).
 3(S)-{2-[5-Benzyl-2-oxo-3(S)-(3-phenylpropionylamino)2,3,4,5-tetrahydrobenzo[b][1,4]diazepin-1-yl]acetylamino)-4-oxobutyric acid tert-butyl ester
 semicarbazone was deprotected as described in the
- 15 preparation of compound K (step C) to give 168 mg (81%) of the title compound as a white solid.

 18 NMR (CD₃OD)

 5 7.37-7.0 (14H, m), 4.75-4.62 (1H, m), 4.6-4.45 (2H, m), 4.4-4.21 (2H, m), 4.15-3.95 (2H, m), 3.15-3.0 (2H, m), 2.82-2.67 (2H, m), 2.65-2.52 (1H, m), 2.5-2.32 (3H, 20 m).

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2,6-Dichlorobenzoic acid 4-tert-butoxycarbonyl-2-oxo-3(s)-(2-[2-oxo-5-(3-phenylpropionyl)-3(s)-(3-phenylpropionyl)-3(s)-(3-phenylpropionylamino)-2,3,4,5-tetrahydro-benzo[b][1,4]diazepin-1-yl]acetyl-amino]butyl ester

5 (107a). The resulting semicarbazone was prepared by the coupling of compound 105b and t-butyl 3-(allyloxycarbonylamino)-4-oxo-5-(2,6-dichlorobenzoyloxy)pentanoate (WO 93 16710) as described in compound 56a to give 256 mg (58%) of the title compound as a

10 white solid. ¹H NMR (CDCl₃) δ 7.45-7.04 (17H, m), 6.45-6.34 (2H, m), 5.28-5.21 (1H, m), 5.1-5.0 (1H, m), 4.95-4.90 (1H, m), 4.75-4.70 (1H, m), 4.55-4.44 (1H, m), 4.32-4.22 (1H, dd), 3.99-3.85 (1H, dd), 3.85-3.76 (1H, m), 3.06-2.83 (5H, m), 2.83-2.74 (1H, m), 2.6-2.44 (2H, 15 m), 2.43-2.33 (1H, m), 2.24-2.15 (1H, m), 1.45 (9H, s).

2,6-Dichlorobenzoic acid 4-carboxy-2-oxo-3(S)-(2-[2-oxo-5-(3-phenylpropionyl)-3(S)-(3-phenylpropionyl)-3(S)-(3-phenylpropionylamino)-2,3,4,5-tetrahydrobenzo[b][1,4]diazepin-1-yl]acetylamino]butyl ester (108a) was prepared from 107a by the method described for compound 57a which afforded 156 mg (68%) of the title compound as a white solid.

1 NMR (CD30D) 8 7.5-6.9 (17H, m), 5.16-5.02 (1H, dd), 4.88-4.71 (2H, m), 4.62-4.44 (2H, m), 4.42-4.28 (2H, m), 4.27-4.18 (2H, m), 3.47-3.41 (1H, m), 2.90-2.60 (5H, m), 2.46-2.4 (2H, m), 2.39-2.18 (2H, m).

 $\label{lem:condition} $$4-(7-Methoxybenzoxazol-2-yl)-4-oxo-3(S)-\{2-\{2-oxo-5-(3-phenylpropionyl)-3(S)-(3-phenylpropionylamino)-2,3,4,5-tetrahydrobenzo[b][1,4]diazepin-1-yl]-acetylamino) $$$

30 butyric acid (108b) was prepared by the method

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described for compound **69a** to give the title compound (50%) as a white solid. ^{1}H NMR (CD₃OD) δ 7.41-6.88 (17H, m), 5.6-5.55 (0.5H, t), 5.48-5.43 (0.5H, t), 4.64-4.45 (2H, m), 4.45-4.30 (1H, m), 3.93 (1.5H, s), 5.3.90 (1.5H, s), 3.47-3.34 (1H, m), 3.10-2.85 (2H, m), 2.84-2.63 (5H, m), 2.6-2.4 (2H, m), 2.3-2.1 (2H, m).

t-Butyl (3S) N-(allyloxycarbonyl)-3-amino-5-(2-chlorophenylmethylthio)-4-oxo-pentanoate (123).

Potassium fluoride (273mg, 4.70mmol) and then 210 chlorophenylmethyl thiol (373mg, 2.35mmol) were added
to a stirred solution of (35) t-butyl N(allyloxycarbonyl)-3-amino-5-bromo-4-oxo-pentanoate
(122; 749mg, 2.14mmol; WO 93 16710) in
dimethylformamide (20ml). The mixture was stirred for
15 3.5h, quenched with water (50ml) and extracted with
ethyl acetate (2 x 50ml). The combined organic
extracts were washed with water (4 x 50ml) then prine

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(50ml). They were dried (MgSO₄) and concentrated to afford an oil which was purified by flash chromatography (10-35% ethyl acetate/hexane) to afford 832 mg (91%) of a colourless solid: mp. 45-6 °C;[α] $_D$ ²⁰ -19.0° (c 1.0, CH₂Cl₂); IR (film) 3340, 2980, 2985, 1275, 1273, 1273, 1274, 1474, 1474, 1474, 1475,

1725, 1712, 1511, 1503, 1474, 1446, 1421, 1393, 1368, 1261, 1244, 1157, 1052, 1040, 995, 764, 739; [↑]H NMR (CDCl₃) δ 7.36 (2H, m), 7.21 (2H, m), 5.97 (2H, m), 4.76 (1H, m), 4.59 (2H, d), 3.78 (2H, s), 3.36

10 (2H, m), 2.91 (1H, dd), 2.74 (1H, dd), 1.43 (9H, s).
Anal. Calcd for C₂₀H₂₆ClNO₅S: C, 56.13; H, 6.12; N,
3.27; S, 7.49. Found: C, 56.08; H, 6.11; N, 3.26; S,
7.54. MS (C.I.) 430/28 (M* + 1, 3%), 374/2 (100).

t-Butyl (3S) 3(2(6-benzyl-1,2-dihydro-2-oxo-3(3-

- 15 phenylpropionylamino)-1-pyridyl)acetylamino-5-(2chlorophenylmethylthio)-4-oxopentanoate (124a). 6Benzyl-1,2-dihydro-2-oxo-3-(3-phenylpropionylamino)pyridyl acetic acid (52b; 300mg, 0.76mmol) in THF (7ml)
 was stirred with 1-hydroxybenzotriazole (205mg,
- 20 1.52mmol) and 1-(3-dimethylaminopropy-3ethylcarbodiimide hydrochloride). After 3 min, water (12 drops) was added and the mixture stirred 10min then treated with t-butyl (35) N-(allyloxycarbonyl)-3-amino-5-(2-chlorophenylmethylthio)-4-oxopentanoate (123)
- 25 (325mg, 0.76mmcl), bis (triphenylphosphine) palladium II chloride (20mg) and tributyltin hydride (0.6mt, 2.28mmol). The mixture was stirred for 5h at room temperature, poured into cthyl acetate and washed with aqueous IM HCl (x2), aqueous sodium bicarbonate, brine,
- 30 dried (MgSO₄) and concentrated. The residue was triturated with pentane and the supernatant discarded. Chromatography (silica gel, 50% ethyl acetate/hexane)

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afforded a colourless foam (439mg, 81%): $\{\alpha\}_D^{21} - 18.3^{\circ}$ (c 0.5, $\mathrm{CH_2Cl_2}$); IR (KBr) 3356, 3311, 1722, 1689, 1646, 1599, 1567, 1513, 1367, 1154; $^1\mathrm{H}$ NMR (CDCl_3) δ 8.39 (1H, d), 8.23 (1H, s), 7.24 (14H, m), 6.16 (1H, d), 4.95 (1H, m), 4.63 (2H, m), 4.02 (2H, s), 3.74 (2H, s), 3.27 (2H, s), 2.85 (6H, m), 1.40 (9H, s). Anal. Calcd for $\mathrm{C3gH_4ClN_3O_6S}$: C, 65.39; H, 5.91; N, 5.87. Found: C, 65.51; H, 5.99; N,5.77.

t-Butyl [3S(1S,9S)]-3-(6,10-dioxo-1,2,3,4,7,8,9,10-10 octahydro) -9-(3-phenylpropionylamino) -6Hpyridazine[1,2-a][1,2]diazepine-1-carboxamido-5-(2chlorophenylmethylthio)-4-oxopentanoate (124b) was prepared by a similar method as 124a from the thioether 123 and 3S(1S, 9S) - 3 - (6, 10 - diox - 1, 2, 3, 4, 7, 8, 9, 10 - 1)15 octahydro)-9-(3-phenylpropionylamino)-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxylic acid (45a) to afford 452mg (50%) of colourless foam: mp 55-7 °C; $[\alpha]_{D}^{22}$ -94.0° (c 0.12, CH₂Cl₂); IR (KBr) 3288, 2934, 1741, 1722, 1686, 1666, 1644, 1523, 1433, 1260, 1225. 20 1146, 757; ¹H NMR (CDCl₃) δ 7.35 (3H, m), 7.20 (7H, m), 6.46 (1H, d), 5.21 (1H, m), 4.97 (2H, m), 4.56 (1H, m), 3.75 (2H, s), 3.25 (3H, m), 2.93 (5H, m), 2.71 (1H, dd), 2.55 (2H, m), 2.30 (1H, m), 1.92 (3H, m), 1.66 (2H, m), 1.42 (9H, s). Anal. Calcd for C35H43ClNaO7S. 25 0.25H₂O: C, 59.73; H, 6.23; Cl, 5.04; N, 7.96; S. 4.56. Found: C, 59.73; H, 6.19; Cl, 5.10; N, 7.79; S, 4.58. MS (-FAB) 697 (M-1, 100).

(3S) 3(2(6-Benzyl-1,2-dihydro-2-oxo-3-(3-phenylpropionylamino)-1-pyridyl)acetylamino-5-(2-30 chlorophenylmethylthio)-4-oxopentanoic acid (125a).

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t-Butvl-3(2(6-benzvl-1,2-dihydro-2-oxo-3-(3phenylpropionylamino)-1-pyridyl)acetyl-amino-5-(2chlorophenylmethylthio) -4-oxopentanoate (124a) (400mg, 0.56mmol) in dichloromethane (3ml) at 0 °C was treated 5 with trifluoroacetic acid (3ml) and stirred at 0 °C for 1h and room temperature for 0.5h. The solution was concentrated then redissolved in dichloromethane and reconcentrated. This procedure was repeated three times. The residue was stirred in ether for 1hr and 10 filtered to yield a colourless solid (364mg, 99%): mp. 165-7 °C; [α]η²² -27.7 ° (c 0.2, CH₂Cl₂); IR (KBr) 3289, 1712, 1682, 1657, 1645, 1593, 1562, 1527, 1497, 1416, 1203, 1182; ¹H NMR (CDCl₃) d 8.47 (1H, d), 8.21 (1H, s), 7.70 (1H, d), 7.22 (14H, m), 6.24 (1H, d), 5.03 15 (1H, m), 4.65 (2H, m), 4.06 (2H, s), 3.69 (2H, m), 3.23 (2H, m), 2.88 (6H, m).

[3S(1S,9S)]-3-(6,10-dioxo-1,2,3,4,7,8,9,10-octahydro)-9-(3-phenylpropionyl-amino)-6H-pyridazine[1,2-a][1,2]diazepine-1-carboxamido-5-(2-

- 20 chlorophenyl-methylthio)-4-oxopentanoic acid (125b),
 was prepared by a similar method as 125a from the t butyl ester 124b to afford 362mg (93%) of colourless
 powder: mp 76-80 °C; [α]_D²¹ -134 ° (c 0.10, MeOH); IR
 (KBr) 3309, 2935, 1725, 1658, 1528, 1445, 1417, 1277,
- 25 12'9, 1175; ^{1}H NMR (D₆-DMSO) δ 8.80 (1H, d), 8.19 (1H, d) 7.31 (9H, m), 5.09 (1H, m), 4.74 (1H, m), 4.63 (1H, m), 4.35 (1H, m), 3.76 (2H, m), 3.28 (3H, m), 2.80 (5H, m), 2.52 (4H, m), 2.16 (2H, m), 1.90 (3H, m). Anal. Calcd for $C_{31}H_{35}Cl_{2}N_{4}O_{7}S$. 0.25H₂O: C, 57.49; H, 5.53;
- 30 N, 8.65; S, 4.95. Found: C, 57.35; H, 5.43; N, 8.45; S, 4.88. MS (-FAB) 641 (M-1, 100).

2-Chlorophenylmethyliodide. A mixture of 2-chlorophenylmethylbronide (4g, 19.47mmol) and NaI (14g, 97.33mmol) in acetone (40ml) was stirred under reflux for 1 hour. The reaction mixture was cooled, filtered and concentrated in vacuo. The residue was triturated with hexane and filtered. The solution was concentrated in vacuo, and the resulting oil purified by flash chromatography (silica, hexane) to afford the title compound (4.67g, 63%) as an oil: ¹H NMR (CDC13)

(3S) t-Butyl N-(allyloxycarbonyl)-3-amino-5-(2-chlorophenylmethyloxy)-4-oxopentanoate (201). (3S) t-Butyl N-(allyloxycarbonyl)-3-amino-5-hydroxy-4-oxopentanoate (81, Chapman, et al., <u>Bioorg. & Med.</u>

15 <u>Chem. Lett.</u>, 2, pp. 613-618 (1992) 0.144g, 0.5mmol) and

10 8 7.34 (4H, m), 4.54 (2H, s).

- 2-chlorophenylmethyliodide (0.569g, 1.5mmol) in CH₂Cl₂ (4ml) were stirred vigorously with silver oxide (0.231g, 1mmol) and heated at 38 °C for 40 hours. The reaction mixture was cooled, filtered and the filtrate
- evaporated. The residue was purified by flash chromatography (silica, 0-20, ethylacetate in hexano) to afford the product as a colourless oil (0.136g, 67%): $\left(\alpha\right)_{D}^{24} + 3.9$ ° (c 1.3, CH₂Cl₂); 1 H NMR (CDCl₃) & 7.37 (4H, m), 5.88 (2H, m), 5.26 (2H, m), 4.69 (2H, s),
- 25 4.57 (3H, m), 4.50 (1H, d), 4.35 (1H, d), 3.03 (1H, dd), 2.76 (1H, dd), 1.42 (9H, s).

- 5,7-Dichlorobenzoxazole (203). A solution of 2,4-dichloro-6-nitrophenol (202, 40g containing 20% moisture) in EtOAc (500ml) was dried using MgSO₄, filtered and the filter cake washed with a little
- 5 EtOAc. Platinum on carbon (5% sulphided 2g) was added and the mixture hydrogenated until uptake of H₂ ceased. Triethyl orthoformate (160ml) and p-toluene sulphonic acid (160mg) were added and the mixture refluxed for 4h. After cooling and removal of spent
- 10 catalyst by filtration the solution was washed with sat. NaHCO₃ solution, water and brine, dried with MgSO₄ and evaporated to dryness. Trituration with hexane gave a solid which was collected by filtration, washed with hexane and dried to give the title compound
- 15 (25.5g, 88%) as a crystalline solid: mp 98-99 °C; IR (KBr) 3119, 1610, 1590, 1510, 1452, 1393, 1296, 1067, 850; $^{1}{\rm H}$ NMR (CDCl₃) δ 8.16 (1H, s), 7.69 (1H, d, J = 1.9), 7.42 (1H, d, J = 1.9); Anal. Calcd for C₇H₃Cl₂NC: C, 44.72; H, 1.61; N, 7.45; Cl, 37.70. Found: C,
- 20 44.84; H, 1.69; N, 7.31; Cl, 37.71.

(3S,4RS) t-Butyl N-(allyloxycarbonyl)-3-amino-4hydroxy-4-(5,7-dichlorobenzoxazol-2-yl)butanoate (204). Magnesium bromide was prepared by reaction of Mg (7.45g, 0.30mole) in THF (516ml) with 1₂ (50mg) and 25 1,2-dibromoethane (26.3ml, 57.3g, 0.30mole) at reflux

for 2h and then cooling to -40 °C. To the above was

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added rapidly Via cannula a solution of 2-lithio-5,7-dichlorobenzoxazole at 70 °C (prepared from 5,7-dichlorobenzoxazole (203, 28.9g, 0.154mole) and butyl lithium (100ml 1.52M in hexane) in THF (150ml) at -

- 5 70 °C). The mixture was stirred at -40 °C for 1h and then cooled to -70 °C before adding a solution of (35) t-butyl N-(allyloxycarbonyl)-3-amino-4-oxo-butanoate (Chapman, et al., Bioorg. 6 Med. Chem. Lett., 2, pp. 613-618 (1992)) (20.3g, 0.078mole) in THF (160ml) at
- 10 less than -60 °C. The reaction was allowed to warm to ambient temperature and was stirred for 16h before quenching with ammonium chloride solution and extracting with 1:1 hexane:ethylacetate 600ml. The organic solution was washed with water and brine, dried
- with MgSO₄ and evaporated to a syrup (52.9g). Flash chromatography (SiO₂ 250g -11 aliquots of 1:1 hexane: CH₂Cl₂ x2, CH₂Cl₂, 5% EtOAc in CH₂Cl₂, 10% EtOAc in CH₂Cl₂, 20% EtOAc in CH₂Cl₂) gave impure product 24.6g which on further chromatography (SiO₂ 1:1 hexane:ether)
- 20 give the title compound as a golden-brown glass (22.7g, 64%); IR (film) 3343, 2980, 1723, 1712, 1520, 1456, 1398, 1369, 1254, 1158, 993; ¹H NMR (CDCl₃) & 7.60 (1H, m), 7.37 (1H, m), 5.72 (1H, m), 5.64 (0.5H, d), 5.10 (2.5H, m), 4.7-4.3 (4H, m), 2.9-2.6 (2H, m), 1.46 and
- 25 1.42 (9H combined, 2 x s). MS ES* Da/e 445 (M + 1)* Cl 35 62%, 447 (M + 1)* Cl 37 40%, 389 100%.

) 1 H NMR (D₆-DMSO) δ 6.10 (1H, d), 5.96-5.88 (1H, m), 5.31-5.12 (2H, m), 4.45 (2H, m), 3.90-3.84 (1H, t), 2.18 (2H, m), 1.85-1.76 (2H, m), 1.36 (9H, s).

(2R)-N-Allyloxycarbonyl-5-(1,1-dimethylethyl)glutamate (205b), was prepared by an analogous method to 205a to 15 afford a colourless oil (6.27g, 88%): [α]_D²⁰ +16° (c 0.095, MeOH); IR (KBr) 3678, 3332, 3088, 2980, 2937, 1724, 1530, 1453, 1393, 1370, 1331, 1255, 1155, 1056, 995. 935. 845, 778, 757, 636, 583; ¹H NMR (CDCl₂) 5

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9.24 (1H, broad s), 5.94-5.79 (1H, m), 5.58 (1H, d), 5.33-5.17 (2H, m), 4.55 (2H, d), 4.38-4.31 (1H, m), 2.41-1.95 (4H, m), 1.42 (9H, s); Anal. Calcd for C₁₃H₂₁NO₆: C, 54.35; H, 7.37; N, 4.88. Found: C, 54.4; H, 7.5; N, 4.8.

(4S) t-Butvl N-allyloxycarbonyl-4-amino-5-

hydroxypentanoate (206a). To a solution of 205a (3.6g, 12.5mmol) in THF (100ml) at 0 °C was added N-methyl morpholine (1.5ml, 13mmol) followed by isobutyl

- 10 chloroformate, (1.1ml, 13mmol). After 15 minutes, this mixture was added to a suspension of NaBH₄ (0.95g, 25mmol) in THF (100ml) and MeOH (25ml) at -78 °C. After 2 hours at -70 °C, the mixture was quenched with acetic acid, diluted with EtOAc, washed with a sat.
- 15 hydrogenocarbonate solution 3 times, water and a sat. solution of salt, dried and evaporated. Flash chromatography (2% MeOH in $\mathrm{CH_2Cl_2}$) afforded 206a as a colourless oil (2.4g, 70%): $[\alpha]_D^{20}$ -10 ° (c 3.8%, $\mathrm{CH_2Cl_2}$) ¹H NMR (CDCl₃) & 5.84 (1H, m), 5.34-5.17 (3H, 20 m), 4.56-4.53 (2H, m), 3.68-3.59 (2H, m), 2.98 (1H, m, 2.40-2.30 (2H, t), 1.84-1.78 (2H, m), 1.43 (9H, s):
- 2.40-2.30 (2H, t), 1.84-1.78 (2H, m), 1.43 (9H, s); Anal. Calcd for C₁₃H₂₃NO₅: C, 57.13; H, 8.48; N, 5.12. Found: C, 57.1; H, 8.6; N, 6.0

(4R) t-Butyl N-allyloxycarbonyl-4-amino-5-

- 25 hydroxypentanoate (206b), was prepared by an analogous
 method to 206a which afforded the title compound as a
 light yellow oil (3.42g, 57%): [ac.] 20 +14 (c 0.166,
 MeOH); IR (KBr) 3341, 3083, 2976, 2936, 2860, 1724,
 1533, 1454, 1419, 1369, 1332, 1251, 1156, 1062, 997,
 30 933, 846, 777, 647; ¹H NMR (CDCl₃) & 5.98-5.81 (1H, m.)
 - 5.35-5.10 (3H, m), 4.55 (2H, d), 3.70-3.56 (3H, m),

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2.50-2.47 (1H, broad s), 2.37-2.30 (2H, π), 1.89-1.74 (2H, m), 1.44 (9H, s); Anal. Calcd for $C_{13}H_{23}NO_5$: C, 57.13; H, 8.48; N, 5.12. Found: C, 56.9; H, 8.6; N, 5.6

- 5 (4S) t-Butyl N-Allyloxycarbonyl-4-amino-5-oxopentanoate (207a). To a solution of DMSO (1.51g, 19.3mmol) in CH₂Cl₂ (25ml) at -70 °C was added oxalyl chloride (1.34g, 19.3mmol). After 10 minutes at -70 °C, a solution of (206a) (2.4g, 8.8mmol) in CH₂Cl₂ (10ml) was added dropwise and the mixture stirred for 15 minutes at -70 °C. Diisopropylethylamine (3.4g, 26.3mmol) was added and the mixture stirred at -25 °C for 15 minutes then diluting with EtOAc (50ml) washed with a solution of sodium hydrogen sulfate 2M, concentrated to give an 15 oil which was used immediately without purification:

 1 NMR (CDCl₃) δ 9.5 (1H, s), 6.0-5.5 (2H, m), 5.5-5.1 (2H, m), 4.5 (2H, m), 4.2 (1H, m), 2.4-2.10 (2H, m), 2.05 (2H, m), 1.36 (9H, s).
- (4R) t-Butyl N-Allyloxycarbonyl-4-amino-5-oxopentanoate 20 (207b), was prepared by an analogous method to 207a which afforded an oil (2.95g, 96%) which was used without further purification in the next step: $\left[\alpha\right]_{\rm D}^{20}$ +21 ° (c 0.942, MeOH); 1 H NMR (CDCl $_{3}$) δ 9.58 (1H, s), 6.05-5.80 (1H, m), 5.57 (1H, broad s), 5.35-5.18 (2H, 25 m), 4.56 (2H, d), 4.34-4.24 (1H, m), 2.38-2.16 (3H, m), 1.96-1.73 (1H, m), 1.43 (9H, s).
- (4S) t-Butyl N-Allyloxycarbonyl-4-amino-5-oxopentanoate
 semicarbazone (208a). To a solution of 207a (2.39g,
 8.8mmol), in MeOH (20ml) was added sodium acetate
 30 (0.72q, 8.8mmol) and semicarbazide (0.98q, 8.8mmol)

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stirred overnight, concentrated and diluted with CH₂Cl₂ (100ml), washed with water, dried and concentrated. Flash chromatography (2% MeOH in CH₂Cl₂) afforded 208a (2.10g, 73%) as an oil: $\{\alpha\}_{\mathbf{D}}^{20}$ -21 (c 2.55 °, CH₂Cl₂); 5 1 H NNR (CDCl₃) δ 9.98 (1H, s), 7.27 (1H, d), 5.8 (1H, m), 5.5 (1H, d), 5.35-5.19 (2H, m), 4.58 (2H, m), 4.14 (1H, m), 2.37 (2E, t), 2.09 (1H, m), 2.0-1.75 (2H, m); Anal. Calcd for C₁₄H₂₄N₄O₅: C, 51.21; H, 7.37; N, 17.06. Found: C, 50.2; H, 7.3; N, 16.1

10 (4R) t-Butyl N-Allyloxycarbonyl-4-amino-5-oxopentanoate semicarbazone (208b), was prepared by an analogous method to 208a which afforded a glassy oil (2.37g, 66%): [α]_D²⁰ +30 (c 0.26, CHCl₃); IR (KBr) 3476, 3360, 2979, 2923, 1700, 1586, 1527, 1427, 1394, 1369, 1338, 15 1253, 1156, 1060, 997, 929, 846, 775; ¹H NMR (CDCl₃) δ 9.87 (1H, s), 7.09 (1H, d), 6.05-5.75 (3H, m), 5.58 (1H, d), 5.32-5.16 (2H, m), 4.54 (2H, d), 4.35 (1H, m), 2.32-2.26 (2H, m), 2.15-1.55 (2H, m), 1.41 (9H, s);

Anal. Calcd for $C_{14}H_{24}N_{4}O_{5}$: C, 51.21; H, 7.37; N,

20 17.06. Found: C, 51.0; H, 7.5; N, 16.7.

211 (b)
$$R^1 = MeSO_2$$

(c)
$$R^1 = MeCO$$

(d)
$$R^1 = PhCH_2OCO$$

(e)
$$R^1 = PhCO$$

$$(f) R^1 = Fmoc$$

212 (b)
$$R^1 = Meso_{\sim}$$

(c)
$$R^1 = MeCO$$

(d)
$$R^1 = PhCH_0OCO$$

(e)
$$R^1 = PhC0$$

$$(f) R^1 = Fmoc$$

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- (15,95) t-Butyl 6,10-dioxo-9-methylsulphonylamino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxylate (211b). A solution of t-butyl 9-amino-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2]diazonylo-1
- 5 octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxylate (GB 2,128,984; 831mg, 2.79mmol) and disopropylethylamine (1.22ml, 6.99mmol, 2.5 equiv) in CH₂Cl₂ (10ml) under dry nitrogen was treated with methanesulphonyl chloride (237pl, 3.07mmol 1.1 equiv).
- 10 The mixture was stirred for 1h, diluted with EtOAc (75ml) and washed with saturated NaHCO $_3$ (50ml) and saturated aqueous sodium chloride (30ml), dried (MgSO_4) and concentrated. Flash chromatography (10--35% EtoAc in CH $_2$ Cl $_2$) afforded 211b (806mg, 77%) as a colourless
- 15 solid: mp 68-70 °C; $[\alpha]_D^{23}$ -109 (c 1.09, CH_2Cl_2); IR (KBr) 3270, 2980, 2939, 1735, 1677, 1458, 1447, 1418, 1396, 1370, 1328, 1272, 1252, 1232, 1222, 1156, 1131, 991; 1H NMR (CDCl₃) δ 6.15 (1H, d), 5.31 (1H, m), 4.65-4.11 (2H, m), 3.47 (1H, m) 2.99 (3H, s), 2.89 (1H,
- 20 m), 2.72-2.51 (2H, m), 2.34 (1H, m), 2.26 (1H, m;, 2.05-1.62 (4H, m), 1.47 (9H, s); Anal. Calcd for $C_{15}H_{23}N_{3}O_{6}S$: C, 47.97; H, 6.71; N, 11.19; S, 8.54. Found: C, 48.28; H, 6.68; N, 10.86; S, 8.28. MS (+FAB) 376 (M* + 1, 66%), 320 (100).
- 25 (15,95) t-Butyl 9-acetylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino [1,2-a][1,2]diazepine-1-carboxylate (211c). Acetic annyoride
 (307mg, 3.01mmol) was added to a stirred mixture of tbutyl 9-amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-
- 30 pyridazino[1,2-a][1,2]diazepine-1-carboxylate
 (GB 2,128,984; 813.7mg, 2.74mmol),
 disopropylethylamine (884mg, 6.84mmol) and CH₂Cl₂

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(20ml). The mixture was kept for 1h then diluted with EtOAc, washed with NaHCO3 solution then brine, dried (MgSO4) and concentrated to yield a colourless oil. The product was purified by flash chromatography (0.5-8% MeoH/CH2Cl2) to afford 211c (804mg, 71%) of colourless powder: mp 162-3 °C; [c]p^23 -109 (c 1.03, CH2Cl2); IR(KBr) 3358, 2974, 1733, 1693, 1668, 1528, 1462, 1431, 1406, 1371, 1278, 1271, 1250, 1233, 1217, 1154, 1124; δ 1 H NMR (CDCl3) d 6.32 (1H, d), 5.29-5.25 10 (1H, m), 4.98-4.85 (1H, m), 4.68-4.58 (1H, m), 3.55-3.39 (1H, m), 2.91-2.66 (2H, m), 2.39-2.18 (2H, m), 2.03 (3H, s), 1.88-1.64 (4H, m), 1.47 (9H, s); Anal. Calcd for $C_{16}H_{25}N_{3}O_{5}$: C, 56.62; H, 7.43; N, 12.38.

Found: C, 56.62; H, 7.43; N,12.36; MS (+ FAB) 340 (M⁺ 15 + 1, 40%), 284 (100).

(15,95) t-Butyl 9-(benzyloxycarbonylamino)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2] diazepine-1-carboxylate (211d). Benzyl chloroformate (1.07g) was added dropwise to a stirred ice cold

- 20 mixture of the (1S,9S) t-butyl 9-amino-6,10-dloxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a]
 [1,2]diazepine-i-carboxylate (GB 2,128,984; 1.55g,
 5.2lmmol), NaHCO₃ (0.66g, 7.82mmol), dioxan (32ml) and
 water (8ml). The mixture was kept at 5 °C for 15min
- 25 then for 2h at room temperature. The mixture was diluted with EtOAc (50ml), washed twice with sat. ${\tt NaHCO_3} \ \, {\tt Solution}, \ \, {\tt dried} \ \, ({\tt MgSO_4}) \ \, {\tt and} \ \, {\tt concentrated}. \ \, {\tt The} \ \, {\tt oily} \ \, {\tt residue} \ \, {\tt was} \ \, {\tt purified} \ \, {\tt by} \ \, {\tt flash} \ \, {\tt chromatography} \ \, {\tt to} \ \, {\tt afford} \ \, {\tt 211d} \ \, (1.98g, 86\$) \ \, {\tt of} \ \, {\tt a colourless oil:} \ \, {\tt [\alpha]}_{\tt D}^{24} \ \, -$
- 30 56.4 (c 1.0, CH₂Cl₂); IR(thin film) 3325, 2979, 2946, 1728, 1677, 1528, 1456, 1422, 1370, 1340, 1272, 1245, 1156, 1122, 1056, 916, 734, 699; ¹H NMR (CDCl₂) 8 7.29

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 $(5H, m), 5.81-5.72 (1H, m), 5.26-5.20 (1H, m), 5.05 \\ (2H, s), 4.69-4.51 (2H, m), 3.48-3.36 (1H, m), 2.81-2.51 (2H, m), 2.34-2.19 (2H, m), 1.90-1.54 (4H, m), 1.41 (9H, s); Anal. Calcd for <math>C_{22}H_{29}N_3O_6^{*}H_2O$: C, 58.79; 5 H, 6.92; N, 9.35. Found: C, 59.10; H, 6.57; N, 9.25; MS (ES +) 454 (M*+Na, 878), 432 (M*+1, 100).

(1S,9S) t-Butyl 9-benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxylate (211e). A solution of benzovl 10 chloride (1.61g, 11.47mmol) in CH2Cl2 (15ml) was added dropwise to a stirred ice cold mixture of (15,95) tbutyl 9-amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxylate (GB 2,128,984; 3.1g, 10.43mmol), dry $\mathrm{CH_2Cl_2}$ (20ml) and 15 diisopropylethylamine (4.54ml, 26.06mmol). The mixture was kept cold for 1h then left at room temperature for 0.5h. The mixture was diluted with CHoClo, washed twice with brine, dried (MgSO4) and concentrated. The residue was purified by flash chromatography (0-5% 20 metha: 1 in CH₂Cl₂) to afford **211e** (4.0q, 96%) of a colourless glass: mp 74-76 °C; $[\alpha]_{D}^{3C}$ -75.0 ° (c 0.12, CH₂Cl₂). IR (KBr) 3350, 2979, 2938, 1736, 1677, 1662, 1536, 1422, 1276, 1250, 1155; ¹H NMR (CDCl₃) δ 8.72 (2H, m), 7.53-7.40 (3H, m), 7.07 (1H, d, J = 7.2), 5.3025 (1H, dd, J = 3.0, 5.8), 5.12 (1H, m), 4.66 (1H, m), 3.51 (1H, m), 2.90 (2H, m), 2.38 (1H, dd, J 13.2, 6.8). 2.25 (1H, m), 1.9 (2H, m), 1.70 (1H, m). Anal. Calcd

36 (1S,9S) t-Butyl 6,10-dioxo-9-(fluoren-9-ylmethyloxy-carbonylamino)-1,2,3,4,7,8,9,10-octahydro-6H-

Found C, 61.69; H, 6.71; N, 10.18.

for Co1Ho2N3Os 0.5H2O: C, 61.45; H, 6.88; N, 10.24.

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pyridazino[1,2-a][1,2]-diazepine-1-carboxylate (211f), was prepared in a similar manner to 211e, except 9-fluorenylmethylchloroformate was used instead of benzoylchloride to give a white glassy solid 211f .

- 5 (2.14g, 898): mp 190-192 °C; $[\alpha]_{\mathbf{D}}^{25}$ -81.5 ° (c 0.1, CH₂Cl₂). IR (KBr) 3335, 2977, 1731, 1678, 1450, 1421, 1246, 1156, 742; ¹H NMR (CDCl₃) δ 7.60 (2H, m), 7.57 (2H, m), 7.50-7.26 (4H, m), 5.60 (1H, d, J = 7.8), 5.28 (1H, m), 4.67 (2H, m), 4.38 (2H, m), 4.23 (1H, m),
- 10 3.59-3.41 (1H, m), 2.92-2.65 (2H, m), 2.41-2.21 (2H, m), 1.95-1.58 (4H, m), 1.47 (9H, s). MS(ES, m/z; 520 (M, + 1, 97%), 179 (100%).

(15,9s) 6,10-Dioxo-9-methysulphonylamino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

- 15 [1,2-a][1,2]diazepine-1-carboxylic acid (212b), was
 synthesized by the same method as compound 212e (635mg,
 85%) as a colourless powder: mp 209-12 °C; [a]_D²⁴ -132
 (c 0.12, MeOH); IR (KBr) 3308, 2940, 1717, 1707, 1699,
 1619, 1469, 1456, 1442, 1417, 1391, 1346, 1339, 1330.
- 20 1310, 1271, 1247, 1222, 1175, 1152, 1133, 993, 976; 2 H NMR (CD₃OD) δ 5.35 (1H, m), 4.58-4.48 (1H, m), 4.46-4.36 (1H, m), 3.60-3.42 (1H, m), 3.01-2.67 (1H, m), 2.95 (3H, s), 2.55-2.39 (1H, m), 2.32-2.26 (2H, m), 2.09-1.89 (2H, m), 1.78-1.62 (2H, m); Anal. Calcd for
- 25 $C_{11}H_{17}N_3O_6S$: C, 41.37; H, 5.37; N, 13.16; S, 10.04. Found: C, 41.59; H, 5.32; N, 12.75; S, 9.76; MS(ES -+: Accurate Mass calculated for C11 $H_{18}N_3O_6S$ (MH *): 320.0916. Found: 320.0943.

(1S,9S) 9-Acetylamino-6,10-dioxo-1,2,3,4,7,8,9,10-

30 octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1carboxylic acid (212c), was prepared from 211e the same

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method as compound 212e as a white glassy solid (595 $_{
m mg}$, 778): mp >250 °C; [α] $_{
m D}^{24}$ -153 (c 0.10, MeOH); IR (KBr) 3280, 2942, 1742, 1697, 1675, 1650, 1616, 1548, 1470, 1443, 1281, 1249, 1202, 1187, 1171; $^{1}_{
m H}$ NMR (CD $_{
m 3}$ CD) δ 5 5.35-5.31 (1H, m), 4.81-4.71 (1H, m), 4.61-4.46 (1H, m), 3.59-3.44 (2H, m), 3.11-2.94 (1H, m), 2.58-2.39 (1H, m), 2.36-2.19 (2H, m), 2.11-1.83 (3H, m), 1.99 (3H, s), 1.78-1.56 (2H, m); Anal. Calcd for C1.9H $_{
m 1}$ N $_{
m 3}$ O $_{
m S}$:

C, 50.88; H, 6.05; N, 14.83. Found: C, 50.82; H,
10 6.02; N, 14.58; MS (ES -) 282 (M-1, 100%): Accurate
 Mass calculated for C₁₂H₁₈N₃O₅ (MH⁺): 284.1246. Found:
284.1258.

(15,95) 9-Benzyloxycarbonylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

- 15 [1,2-a][1,2]diazepine-1-carboxylic acid (212d), was prepared from 211d by the same method as compound 212e as colourless crystals (170mg, 97%): mp 60-100 °C; $\left[\alpha\right]_{D}^{22}$ -103 (c 0.10, MeOH); IR (KBr) 3341, 2947, 1728, 1675, 1531, 1456, 1422, 1339, 1272, 1248, 1221, 1174,
- 20 1122, 1056, 982, 699; ¹H NMR (CDCl₃) & 7.35 (5H, s), 5.65 (1H, d), 5.48-5.40 (1H, m), 5.10 (2H, s), 4.76-4.57 (2H, m), 3.49-3.30 (2H, m), 2.92-2.59 (2H, m), 2.40-2.27 (2H, m), 1.97-1.67 (4H, m); MS (ES -) 374 (M 1, 100%). Accurate mass calculated for C_{1E}H₂₂N₃C₆
- 25 (MH^+) : 376.1509. Found: 376.1483. Accurate mass calculated for $\mathrm{C_{10}H_{21}N_3O_6Na}$ (MNa^+) : 398.1328. Found: 398.1315.

(1S,9S) 9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-

30 carboxylic acid (212e). TFA (20ml) was added to an ice cold stirred solution of the t-butyl ester 211e (4.15g,

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- 10.34mmol) in dry CH_2Cl_2 (20ml). The mixture was kept cold for 1.5h then left for 2.5h at rt, concentrated. TFA was removed by repeated concentrations of CH_2Cl_2 \ether and ether solutions of the residue.
- 5 Finally trituration of the residue with ether afforded
 212e 3.05g (85%) of a white glassy solid: mp 118-
 126 °C; $[\alpha]_D^{24}$ -70.5 ° (c 0.1, CH₂Cl₂). IR (KBr) 3361, 2943, 1737, 1659, 1537, 1426, 1220, 1174;
 ¹H NMR (CDCl₃) δ 7.80 (2H, m), 7.54-7.33 (4H, m), 8.83 (brs),
- 10 5.44 (1H, m), 5.26-5.13 (1H, m), 4.66 (1H, m), 3.59-3.41 (1H, m), 2.97, 2.76 (2H, 2m), 2.36 (2H, m), 1.98 (2H, m), 1.75 (2H, m). MS(ES⁻, m/z) 344 (M - 1, 100%).

(15,95) 6,10-Dioxo-9(fluoren-9-

ylmethyloxycarbonylamino) -1,2,3,4,7,8,9,10-octahydro-15 6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxylic acid

- (212f), was prepared from 211f in 96% yield by the same method as for 212e: mp 120-126 °C; (a)_D²⁵ -72.5 ° (c 0.1, CH₂Cl₂). IR (KBr) 3406, 2950, 1725, 1670, 1526, 1449, 1421, 1272, 1248, 1223, 1175, 761, 741;
- 20 1 H NMR (CDCl₃) δ 7.76 (2H, m), 7.62-7.26 (4H, m), 6.07, 5.76 (2H, brs, d, d, J = 2.9), 5.46, 5.36 (1H, 2m), 4.79-4.54 (2H, m), 4.77 (2H, m), 4.21 (1H, m), 3.41 (1H, m), 2.89 (1H, m), 2.69 (1H, m), 2.35 (2H, m), 1.98, 1.73 (4H, 2m). MS(ES $^{-}$, m/z) 462 (M $^{+}$ 1, 50%),
- 25 240 (100%).

(213) (c) $R^1 = MeCO$ (e) $R^1 = PhCO$

(214) (c) $R^1 = MeCO$ (e) $R^1 = PhCO$

[2RS,3S(1S,9S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-9-(acetylamino)-6,10-dioxo-1,2,3,4,7,8,9,10-

- 5 octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1carboxamide (213c), was synthesized from 212c by the same method as compound 213e to afford a mixture of diastereomers (193mg, 36%) as colourless crystals: IR (KBr) 3272, 1799, 1701, 1682, 1650, 1555, 1424, 1412.
- 10 1278, 1258, 1221, 1122, 937;

 1 M NMR (CDCl₂)

 5 7.41-7.28 (5H, m), 6.52 (0.5H, d), 6.38 (0.5H, d), 6.22 (0.5H, d), 5.57 (0.5H, d), 5.36 (0.5H, s) 5.10-5.05 (1H, m), 5.00-4.45 (5.5H, m), 3.19-2.84 (3H, m), 2.72-2.56 (1H, m), 2.51-2.25 (2H, m), 2.02 (3H, s), 1.98-1.70 (3H, m),
- 15 1.66-1.56 (3H, m); Anal. Calcd for $C_{23}H_{28}N_4O_7$: C, 58.47; H, 5.97; N, 11.86. Found: C, 58.37; H, 6.09; N, 11.47. MS (ES -) 471 (M-1, 100%). Accurate mass calculated for $C_{23}H_{29}N_4O_7$ (MH $^+$): 473.2036. Found: 473.2012. Accurate mass calculated for $C_{23}H_{28}N_4O_7Na$
- 20 (Mna⁺): 495.1856. Found: 495.1853.

[1S,98(2RS,3S)] 9-Benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-N-(2-benzyloxy-5oxotetrahydrofuran-3-yl)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213e). Tributyltin hydride
25 (2.2ml, 8.18mmol) was added dropwise to a solution of

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- acid 212e (1.95g, 5.6mmol), (3s, 2Rs) 3-allyloxycarbonylamino-2-benzyloxy-5-oxotetrahydrofuran (Chapman, Bioorg, 5 Med. Chem. Lett., 2, pp. 615-618 (1992); 1.80g, 6.16mmol) and (Ph₃P)₂PdCl₂ (50mg) in dry
- 5 CH₂Cl₂ (36ml), with stirring, under dry nitrogen.
 After 5 min 1-hydroxybenzotriazole (1.51g, 11.2mmol 6.72mmol) was added followed after cooling (1ce/H₂O) by ethyldimethylaminopropyl carbodiimide hydrochloride (1.29g, 6.72mmol). After 5 mins the cooling bath was
- 10 removed and the mixture was kept at room temperature for 4h, diluted with EtOAc, washed with 1M HCl, brine, sat. aq. NaHCO₃ and brine, dried (MgSO₄) and concentrated. Flash chromatography (silica qcl, 0-90% EtOAc in CH₂Cl₂) gave the product as a white solid
- 15 (2.34g, 78%): IR (KBr) 3499, 1792, 1658, 1536, 1421, 1279, 1257, 1123, 977, 699; ¹H NMR (CDCl₃) δ 7.81 (2H, m), 7.54-7.34 (8H, m), 7.1, 6.97, 6.89, 6.48 (2H, m, d, J 7.7, d, J = 7.5, d, J = 7.6), 5.57, 5.28 (1H, d, J = 5.2, s), 5.23-5.07 (2H, m), 4.93-4.42, 3.22-2.70, 2.51-
- 20 2.26, 2.08-1.69, 1.22 (15H, 5m). Anal. Calcd for $C_{28}H_{30}N_4O_7 \ 0.5H_2O; \ C, \ 61.87; \ H, \ 5.75; \ N, \ 10.32. \ \ Found \\ C, \ 62.02; \ H, \ 5.65; \ N, \ 10.25.$

[3S(1S,9S)] 3-(9-Acetylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

- 25 [1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid
 (214c), was synthesized from 213c by a method similar
 to the method used to synthesize 214e from 213e to
 provide colourless crystals (140mg, 99%): mp 90180 °C; [α]p²² -114 (c 0.10, MeOH); IR (KBr) 3354, 3370,
- 30 2946, 1787, 1658, 1543, 1422, 1277, 1258; $^{1}{\rm H}$ NMR (d $^{6}-$ DMSO) δ 8.66 (1H, m), 8.18 (1H, d), 6.76 (1H, s;, 5.08 (1H, m), 4.68 (1H, m), 4.30 (1H, m), 2.92-2.70 (2H, m),

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2.27-2.06 (3H, m), 1.95-1.72 (4H, m), 1.85 (3H, s), 1.58 (2H, m); MS(ES -) 381 (M-1, 100%); Accurate mass calculated for $C_{16}H_{23}N_4O_7$ (MH $^+$): 383.1567. Found: 383.1548.

5 [3S(1S,9S)] 3-(9-Benzovlamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutancic acid (214e). A mixture of 213e (2.29g, 4.28mmol), 10% palladium on carbon (1.8g) and MeOH (160ml) was stirred under H2 at 10 atmospheric pressure for 6.3h. After filtering and concentrating the hydrogenation was repeated with fresh catalyst (1.8g) for 5h. After filtering and concentrating the residue was triturated with diethyl ether, filtered and washed well with ether to give 214e 15 as a white solid (1.67g, 88%): mp 143-147 °C; $[\alpha a]_{D}^{23}$ -125 ° (c 0.2, CH₃OH). IR (KBr) 3391, 1657, 1651, 1538, 1421, 1280, 1258; ¹H NMR (CD₂OD) δ 7.90 (2H, m), 7.63-7.46 (3H, m), 5.25 (1H, m), 5.08-4.85 (1H, m), 4.68-4.53 (2H, m), 4.33-4.24 (1H, m), 3.62-3.44, 3.22-3.11, 20 2.75-2.21, 2.15-1.92, 1.73-1.66 (11H, 5m). Anal. Calcd for C21H24N4O7 H2O: C, 54.54; H, 5.67; N, 12.11. Found

C, 54.48; H, 5.63; N, 11.92.

- 5 [3s,4Rs(1s,9s)] t-Butyl 3-[9-acetylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido)-5-(2,6-dichlorobenzoyloxy)-4-hydroxypentanoate (215c), was synthesized from 214c by the same method as compound
- 10 215e, to afford a mixture of diastereomers as a white glassy solid (398mg, 84%): IR (KBr) 3336, 2977, 1736, 1658, 1562, 1541, 1433, 1368, 1277, 1150; ¹H NMR (CDCl₃) & 7.36-7.32 (3H, m), 6.91 (1H, d), 6.30 (1H, d), 5.15-5.09 (1H, m) 5.01-4.88 (1H, m), 4.61-4.44 (2H,
- 15 m), 4.37-4.08 (3H, m), 3.32-3.18 (1H, m), 3.04-2.89 (1H, m), 2.82-2.51 (4H, m), 2.39-2.29 (1H, m), 2.08-1.64 (4H, m) 2.02 (3H, s); Anal. Calcd for

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 $\begin{array}{llll} C_{28}H_{34}N_4Cl_2O_9; & \text{C, 52.26; H, 5.64; N, 8.71.} & \text{Found: C,} \\ 52.44; & \text{H, 5.87; N, 8.16.} & \text{MS (ES -) 645/3/1 (M-1, 26%),} \\ 189 & (81), & 134 & (100). & \text{Accurate mass calculated for} \\ & C_{28}H_{37}N_4Cl_2O_9 & (\text{MH}^+); & 643.1938. & \text{Found: 643.1924.} \\ & \text{Accurate mass calculated for $C_{28}H_{36}N_4Cl_2O_9Na (\text{MNa}^+)$} \\ & 665.757. & \text{Found: 665.1756.} \end{array}$

[3S,4RS(1S,9S)] t-Butyl 3-(9-benzyloxycarbonylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(2,6-

10 dichlorobenzyloxy)-4-hydroxypentanoate (215d), was

synthesized from 214d by the same method as compound 215e to afford a mixture of diastereomers (657mg, 70%) as a glassy white solid: IR (KBr) 3420, 3361, 2975, 2931, 1716, 1658, 1529, 1434, 1367, 1348, 1250, 1157,

15 1083, 1055; ¹H NMR (CDCl₃) & 7.32 (8H, m), 7.14 (1H, d), 5.81 (1H, d), 5.15 (1H, m), 5.07 (2H, s), 4.74-4.65 (1H, m), 4.58-4.22 (4H, m), 4.15-4.06 (1H, m), 3.72 (1H, m), 3.32-3.21 (1H, m), 3.04-2.94 (1H, m), 2.69-2.52 (3H, m), 2.33-2.27 (1H, m), 1.95-1.59 (4H, m),

- 20 1.28 (9H, s); Anal. Calcd for $C_{34}H_{40}N_4C1_2O_{10}.0.5\ H_2O$; C, 54.70; H, 5.54; N, 7.50. Found: C, 54.98; H, 5.59; N, 7.24. MS (ES -) 737/5/3 (M-1, 22%), 193/1/89 (100). Accurate mass calculated for $C_{34}H_{41}N_4C1_2O_{10}$ (MH †) 735.2120. Found: 735.2181.
- 25 [3S,4RS(18,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(2,6dichlorobenzyloxy)-4-hydroxypentanoate (215e),
 Tributyltin hydride (4.6ml; 11.4mmol) was added
 30 dropwise to a stirred mixture of (3S,4RS) t-Butyl (Nallyloxycarbonyl)-3-amino-5-(2,6-dichlorobenzoyloxy)-4-

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hydroxypentanoate (prepared by a method similar to the method described in Revesz et al., <u>Tetrahedron. Lett.</u>, 35, pp. 9693-9696 (1994)) (2.64g; 5.7mmol), $(Ph_3P)_2Pdcl_2$ (50mg), CH_2Cl_2 (100ml) and DMF (20ml) at room temperature. The mixture was stirred for a further 10min was then 1-hydroxypenzotriazole (1.54g.

- 5 temperature. The mixture was stirred for a further 10min was then 1-hydroxybenzotriazole (1.54g, 11.4mmol)was added. The mixture was cooled to 0 0 C then ethyldimethylaminopropyl carbodiimide hydrochloride (1.31g; 6.84mmol) added. The mixture was
- 10 kept at this temperature for 15min then at room temperature for 17h. The mixture was diluted with EtOAc (300ml), washed with 1M HCl (2x100ml), sat. aq. NaHCO₃ (3x100ml) and brine (2x100ml), dried (MgSO₄) and concentrated. The residue was purified by flash
- 15 chromatography (2-5% (MeOH/CH₂Cl₂) to afford 3.24g (81%) of 215e as a glassy solid: mp 106-110 °C; IR (KBr) 3354, 1737, 1659, 1531, 1433, 1276, 1150; ¹H NMR (CDCl₃) δ 7.80 (2H, dd, J = 7.9 and 1.5), 7.75-7.26 (6H, m), 7.14-6.76 (2H, m), 5.30-5.02 (2H, m), 4.63-
- 20 4.11 (5H, m), 3.44-3.26 (2H, m), 3.10-2.30 (5H, m), 2.10-1.60 (5H, m), 1.44 (9H, s); Anal. Calcd for $C_{33}H_{38}Cl_2N_4O_9$. 0.75H₂O: C, 55.12; H, 5.54; N, 7.79; Cl, 9.86. Found: C, 55.04; H, 5.34; N, 7.80; Cl, 10.24. MS (ES +) 709/7/5 (M + 1), 378 (59), 324 (64), 322 25 (100).
 - [3s(1s,9s)] t-Butyl 3-(9-acetylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamido)-5-(2,6-dichlorobenzoyloxy)-4-oxopentanoate (216c), was synthesized from 215c by the
- 30 same method as compound **216e** as a glassy white solid (300mg, 83%): mp 80-125 °C; $\left[\alpha\right]_0^{23}$ -89.1 (c 1.08, CH₂Cl₂); IR (KBr) 3356, 2979, 2935, 1740, 1659, 1532,

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1434, 1369, 1276, 1260, 1151; $^1{\rm H}$ NMR (CDCl₃) & 7.39-7.32 (3H, m), 7.13 (1H, d), 6.34 (1H, d), 5.22-5.17 (1H, m), 5.11 (1H, d), 5.04 (1H, d), 4.99-4.88 (2H, m), 4.64-4.52 (1H, m), 3.29-3.11 (1H, m), 3.05-2.67 (4H, 5 m), 2.39-2.29 (1H, m), 2.02 (3H, s), 1.98-1.75 (4H, m), 1.46 (9H, s); Anal. Calcd for $C_{28}H_{34}N_{4}Cl_{2}O_{9}$: C, 52.42; H, 5.34; N, 8.73. Found: C, 52.53; H, 5.70; N, 7.85. MS (ES -) 643/41/39 (M-1, 100%). Accurate mass calculated for $C_{28}H_{35}N_{4}Cl_{2}O_{9}$ (MH $^{+}$): 641.1781. Found: 641.1735. Accurate mass calculated for $C_{28}H_{34}N_{4}Cl_{2}O_{9}Na$ (Mna $^{+}$): 663.1601. Found: 663.1542.

[3s(1s,9s)] t-Butyl 3-(9-benzyloxycarbonylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido)-5-(2,6-

- 15 dichlorobenzoyloxy)-4-oxopentanoate (216d), was
 synthesized from 215d by the same method as compound
 216e to afford 216d as a white glassy solid (688mg,
 68%): mp 90-170 °C; [α]_D²⁵-83.4 (c 1.01, CH₂Cl₂); IR
 (KBr) 3338, 2933, 1736, 1670, 1525, 1433, 1417, 1368,
 20 1258, 1151, 1056, 1031; ¹H NMR (CDCl₂) & 7.33 (8H, m).
- 20 1258, 1151, 1056, 1031; 'H NMR (CDC1₃) & 7.33 (8H, m), 7.18 (1H, d), 5.65 (1H, d), 5.19 (1H, m), 5.09 (2H, s), 4.98-4.86 (1H, m), 4.82-4.49 (2H, d), 3.30-3.07 (1H, m), 3.05-2.59 (4H, m), 2.42-2.27 (1H, m), 2.18-1.59 (5H, m), 1.42 (9H, s); MS (ES-) 737/5/3 (M, 134-, 185 25 (100).

[3s(1s,9s)] t-Butyl 3-(9-benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(2,6dichlorobenzoyloxy)-4-oxopentanoate (216e). Dess30 Martin reagent (3.82g; 9.0mmol) was added to a stirred solution of the alcohol 215e (3.17g; 4.5mmol) in.

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- ${\rm CH_2Cl_2}$ (100ml). The mixture was sirred for 1h, diluted with EtOAc (300ml), then washed with a 1:1 mixture of sat. ${\rm Na_2S_2O_3}$ and sat. ${\rm NaHCO_3}$ (100ml) followed by brine (100ml). The mixture was dried (MgSO₄) then
- 5 concentrated. The residue was purified by flash chromatography to afford 2.2g (70%) of 216e as a colourless solid: mp 102-107 °C; [α]_D³² -92.5 (c 0.1, CH₂Cl₂); IR (KBr) 3374, 2937, 1739, 1661, 1525, 1433, 1275, 1260, 1152; ¹H NMR (CDCl₃) δ 7.85-7.78 (2H, m),
- 10 7.57-7.32 (6H, m), 7.09 (1H, d, J = 7.9), 7.01 (1H, d, J 7.3), 5.25-5.16 (1H, m), 5.16-5.05 (1H, m), 5.15 (1H, d), 5.03 (1H, d), 4.99-4.90 (1H, m), 4.68-4.54 (1H, m), 3.31-3.17 (1H, m), 3.17-2.72 (4H, m), 2.45-2.35 (1H, m), 2.30-1.66 (5H, m), 1.44 (9H, s); Anal. Calcd for
- 15 $C_{33}H_{36}Cl_2N_4O_9$, 0.5 H_2O ; C, 55.62; H, 5.23; N, 7.86; Cl, 9.95. Found: C, 55.79; H, 5.15; N, 7.80; Cl 9.81. MS (ES +) 729/7/5 (M + Na), 707/5/3 (M + 1), 163 (100%).

[3S(1S,9S)] 3-(9-Acetylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

- 20 [1,2-a][1,2]diazepine-1-carboxamido)-5-(2,6-dichlorobenzoyloxy)-4-oxopentanoic acid (217c), was synthesized from 216c by the same method as compound 217e as a glassy white solid (166mg, 66†): mp 85-175 °C; [\alpha]_{0}^{25} -156 (c 0.13, MeOH); IR (KBr) 3373,
- 25 2929, 1742, 1659, 1562, 1533, 1433, 1412, 1274, 1266, 1223, 1197, 1145, 1138; $^1\mathrm{H}$ NMR (CD₃OD) δ 7.38 (3H, s:, 5.14-5.03 (1H, m), 4.49-4.32 (2H, m), 3.50-3.27 (1H, m), 3.11-2.92 (1H, m), 2.84-2.62 (2H, m), 2.46-2.11 (2H, m), 2.05-1.46 (5H, m), 1.92 (3H, s); Anal. Calcd
- 30 for C₂₄H₂₆N₄Cl₂O₉.H₂O: C, 47.77; H, 4.68; N, 9.29. Found: C, 47.75; N, 4.59; N, 9.07. MS (ES +) 627/5/3 (M-K, 215), 611/9/7 (M+Na, 87), 589/7/5 (M⁺ +1, 71),

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266 (100); Accurate mass calculated for $C_{24}H_{27}N_4C1_2O_9$ (MH⁺): 585.1155. Found: 585.1134.

[3S(1S,9S)] 3-(9-Benzyloxycarbonylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-5 diazepine-1-carboxamido)-5-(2,6-dichlorobenzoyloxy)-4-oxopentanoic acid (217d), was synthesized from 216d by the same method as compound 217e to afford 217d as a white glassy solid (310mg, 96%): mp 85-110 °C; [a]_D²⁴-85-9 (c 0.13, MeOH); IR (KBr) 3351, 2945, 1738, 1669, 1524, 1433, 1258, 1147, 1057; ¹H NMR (CD₂OD) 8 7.56 (4H, m), 7.45 (5H, m), 5.32 (2H, m), 5.20 (2H, s), 4.76-4.48 (3H, m), 3.65-3.38 (3H, m), 3.27-3.09 (2H, m), 3.03-2.89 (2H, m), 2.65-2.24 (3H, m), 2.19-1.62 (5H, m); MS (ES -) 679/7/5 (M-1, 100%); Accurate mass 15 calculated for C₃₀H₃₁N₄Cl₂O₁₀ (MH⁺): 677.1417. Found: 677.1430.

[3S(1S,9S)] 3-(9-Benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(2,6-dichlorobenzoyloxy)-4oxopentanoic acid (217e), TFA (25ml) was added dropwise
to an ice cold stirred solution of the ester 216e
(2.11g, 3.0mmol). The mixture was stirred at 0 °C for
20min then at room temperature for 1h. The mixture was
evaporated to dryness then coevaporated with ether

25 three times. Addition of dry ether (50 ml) and filtration afforded 1.9g (98%) of **217e** as a colourless solid: mp 126-130 °C; $(\alpha)_D^{30}$ -122.0 (c 0.1, MeOH); IF (KBr) 3322, 1740, 1658, 1651, 1532, 1433, 1277, 1150; 1 H NMR (D_6 -DMSO) δ 8.87 (HH, d, J = 7.4), 8.61 (HH, d, J = 7.8), 7.92-7.86 (2H, m), 7.65-7.43 (6H, m), 5.25-5.12 (3H, m), 4.94-4.60 (2H, m), 4.44-4.22 (HH, m).

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[3S,4RS(1S,9S)] t-Butyl 4-[5-(2,6-dichlorophenyl)oxazol-2-yl]-3-(6,10-dioxo-9-methylsulphonylamino-10 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

[1,2-a][1,2]diazepine-1-carboxamido)-4-hydroxybutanoate (218b), was prepared from the acid 212b and 99 in an analogous way to compound 215e to afford a mixture of diastereomers (865mg, 80%) as a colourless solid: IR 15 (KBr) 3298, 2974, 1723, 1659, 1544, 1518, 1430, 1394, 1370, 1328, 1273, 1256, 1156, 1134; ¹H NMR (CDCl₃) 5 7.45-7.28 (4H, m), 7.26-7.15 (2H, m), 5.26-5.10 (2H, m), 4.80-4.67 (1H, m), 4.59-4.42 (2H, m), 3.32-3.17

(1H, m), 2.96 (3H, 2xs), 2.93-2.79 (1H, m), 2.71-2.53

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5 636/4/2 (38), 246 (100). Accurate mass calculated for $C_{28}H_{36}N_{5}Cl_{2}O_{9}S$ (MH †): 688.1611. Found: 688.1615.

[3S(1S,9S)]t-Butyl 4-[5-(2,6-dichlorophenyl)-oxazol-2yl]-3-(6,10-dioxo-9-methylsulphonylamino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]di 10 azepine-1-carboxamido)-4-oxobutanoate (219b), was prepared from 218b in an analogous way to compound 216e as an off-white powder (675mg, 81%): mp 100-200 °C; [a]n²⁴ -84.9 (c 1.01, CH₂Cl₂); IR (KBr) 3336, 2978, 2936, 1719, 1674, 1510, 1433, 1421, 1369, 1329, 1274. 15 1257, 1155, 991, 789; ¹H NMR (CDCl₃) δ 7.47-7.38 (4H, m), 7.24 (1H, d), 5.61-5.53 (1H, m), 5.48 (1H, d), 5.38-5.30 (1H, m), 4.67-4.45 (2H, m), 3.48-3.18 (2H, m), 3.04-2.90 (2H, m), 2.97 (3H, s), 2.69-2.54 (1H, m), 2,42-2,32 (1H, m), 2,22-2,15 (1H, m), 2,07-1,93 (3H, 20 m), 1.71-1.65 (2H, m), 1.38 (9H, s); Anal. Calcd for C28H33N3Cl2O9S: C, 48.98; H, 4.84; N, 10.20; S, 4.67. Found: C, 48.73; H, 4.95; N, 9.65; S, 4.54. MS (ES +)

[3S(1S,9S)] 4-[5-(2,6-Dichlorophenyl)oxazol-2-yl]-3-(6,10-dioxo-9-methylsulphonylamino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (220b), was prepared from 219b in an analogous way to compound 217e as a pale cream powder (396mg, 87%): mp 100-200 °C; [0]₁²⁷-

692/90/88 (M $^+$ + 1, 100%), 636/4/2 (71). Accurate mass calculated for $\rm C_{28}H_{34}N_5Cl_2O_9S$ (MH $^+$): 686.1454. Found:

25 686.1474.

129 (c 0.12, MeOH); IR (KBr) 3310, 3153, 1713, 1667, 1557, 1510, 1432, 1421, 1329, 1273, 1258, 1221, 1193, 1153, 1134, 992, 789; 1 H NMR (d⁶ DMSO) δ 7.88 (1H, s), 7.81-7.60 (4H, m), 5.49-5.28 (1H, m), 5.24-5.14 (1H, 5 m), 4.46-4.22 (2H, m), 3.30-3.03 (2H, m), 2.97-2.76 (3H, m), 2.96 (3H, s), 2.46-2.24 (1H, m), 2.16-2.05 (1H, m), 2.03-1.78 (3H, m), 1.68-1.46 (2H, m); MS (ES-) 632/30/28 (M - 1, 688], 149/7/5 (100). Accurate mass

calculated for C24H26N5Cl2OgS (MH+): 630.0828. Found:

10 630.0852.

$$R_1$$
 = $MeSO_2$ 222b R_1 = $MeSO_2$ 222e R_1 $= PhCO$ 223b R_1 = $MeSO_2$ 222e R_1 $= PhCO$ 223b R_1 = $MeSO_2$ 223c R_1 = $MeSO_2$ 223c R_1 = $MeSO_2$ 223e R_1 = $MeSO_2$

15 [3s,4Rs(1s,9s)] t-Butyl 4-(5,7-dichlorobenzoxazol-2yl)-3-(6,10-dioxo-9-methylsulphonylamino1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

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[1,2-a][1,2]diazepine-1-carboxamido)-4-hydroxybutanoate (221b), was prepared from the acid 212b and (35,4RS) t-butyl N-(allyloxycarbonyl)-3-amino-4-hydroxy-4-(5,7-dichlorobenzoxazol-2-yl)butanoate (204) by an analogous method as that used for compound 215e to afford a mixture of diastereomers (460mg, 70%) as a glass: IR (film) 3325, 1725, 1664, 1453, 1399, 1373, 1327, 1274, 1256, 1155; ¹H NMR (CDCl₃) & 7.57 (1H, m), 7.36 (2H, m), 6.06 (1H, t), 5.29 (2H, m), 4.79 (1H, m), 4.47 (1H, 10 m), 3.23 (1H, m), 2.97 and 2.94 (3H combined, 2 x s), 2.9-2.4 (4H, m), 2.30 (1H, m), 1.96 (4H, m), 1.41 and 1.37 (9H combined, 2 x s), MS ES Da/e 660 (M - 1) Cl³⁵ 100%, 662 (M - 1) Cl³⁷.

[3S,4RS(1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo-15 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

[1,2-a][1,2]diazepine-1-carboxamido)-4-(5,7-dichlorobenzoxazol-2-yl)-4-hydroxybutanoate (221e), was prepared from the acid (212e) and (3S,4RS) t-butyl N-(allyloxycarbonyl)-3-amino-4-hydroxy-4-(5,7-

- 20 dichlorobenzoxazol-2-yl)butanoate (204) by an analogous method as that used for compound 215e to afford a mixture of diastereomers (613mg, 87%) as a glass: IR (film) 3328, 1729, 1660, 1534, 1454, 1422, 1399, 1276, 1254, 1155; ¹H NMR (CDCl₃) & 7.80 (2H, d), 7.60-7.35
- 25 (5H, m), 7.05 (2H, m), 5.13 (3H, m), 4.74 (1H, m), 4.51 (1H, m), 3.25 (1H, m), 3.1-2.6 (5H, m), 2.33 (1H, m), 2.1-1.5 (5H, m), 1.43 and 1.41 (9H combined, 2 × s). MS ES⁺ Da/e 688 (M + 1)⁴ Cl^{35} 55%, 690 (M + 1)⁺ Cl^{37} 35%, 328 100%.
- 30 [35(15,95)]t-Butyl 4-(5,7-dichlorobenzoxazol-2-yl)-3-(6,10-dioxo-9-methylsulphonylamino-1,2,3,4,7,8,9,10-

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octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxobutanoate (222b), was prepared from 221b by an analogous method as that used for compound 216e to afford a colourless glass (371mg, 86%): $\left[\alpha\right]_0^{26}$ 5 -81.0 (c 0.1, CH₂Cl₂); IR (KBr) 3324, 2979, 2936, 1726, 1664, 1394, 1370, 1328, 1155, 991; 1 H NMR (CDCl₃) 8 7.78 (1H, d), 7.57 (2H, m), 5.87 (1H, d), 5.69 (1H, m), 5.47 (1H, m), 4.55 (2H, m), 3.24 (2H, m), 3.0 (5H, m + s), 2.59 (1H, m), 2.39 (1H, m), 2.2 - 1.7 (4H, m), 1.65 10 (1H, m), 1.40 (9H, s).

[3S(1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido)-4-(5,7-dichlorobenzoxazo1-2-yl)-4-oxobutanoate (222e), was

15 prepared from 221e by an analogous method as that used

- for compound 216e to afford a colourless glass (480mg, 84%): $(\alpha)_0^{25}$ -86.4 ° (c 0.1 CH₂Cl₂); IR (KBr) 3337, 2978, 2938, 1728, 1657, 1534, 1456, 1422, 1395, 1370, 1277, 1250, 1154; ¹H NMR (CDCl₃) δ 7.80 (3H, m), 7.50 (4H, m), 7.20 (1H, d), 7.02 (1H, d), 5.60 (1H, m), 5.28 (1H, m), 5.15 (1H, m), 4.11 (1H, m), 3.34 (2H, m), 2.96
 - (1H, m), 5.15 (1H, m), 4.11 (1H, m), 3.34 (2H, m), 2.96 (3H, m), 2.40 (1H, m), 2.20 (1H, m), 1.92 (2H, m), 1.67 (2H, m), 1.38 (9H, s). MS ES Da/e 684 (M 1) $^{-}$ C1 35 47%, 686 (M 1) $^{-}$ C1 37 32%.
- 25 [3s(1s,9s)] 4-(5,7-Dichlorobenzoxazo1-2-y1)-3-(6,10dioxo-9-methylsulphonylamino-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1carboxamido)-4-oxobutanoic acid (223b), was prepared
 from 222b by an analogous method as that used for
 30 compound 217e to afford an off-white solid (257mg,
 78%): [α]n²⁵ -105.7 ° (c 0.1, CH₂Cl₂); IR (KBI: 3321,

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1723, 1663, 1407, 1325, 1151, 992; ^1H NMR ($\text{D}_6\text{-DMSO}$) δ 8.96 (1H, d), 8.18 (1H, d), 7.96 (1E, d), 5.50 (1H, m), 5.15 (1H, m), 4.30 (2H, m), 3.06 (2H, m), 2.87 (5H, m + s), 2.29 (1H, m), 1.99 (4H, m), 1.56 (2H, m).

5 [3S(1S,9S)] 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido)-4-(5,7-dichlorobenzoxazol-2-yl)-4-oxobutanoic acid (223e), was prepared from 222e by an analogous method as that used for compound 217e to afford a pale cream solid (311mg, 78%): mp 167-180 °C; [α]_D²³ -88.6 ° (c 0.1 CH₂Cl₂); IR (KBr) 3331, 1724, 1658, 1534, 1458, 1421, 1279, 1256, 991; ¹H NMR (CDCl₃) δ 7.77 (4H, m), 7.4 (5H, m), 5.57 (1H, bs), 5.33 (1H, bs), 5.47 (1H, q), 4.56 (1H, bd), 3.60 (2H, m), 3.20 (3H, m), 2.76 (1H, m), 2.36 (1H, dd), 2.0 (3H, m), 1.66 (1H, m). MS ES Da/e 628 (M - 1)⁻ Cl³⁵ 7%, 630 (M - 1)⁻ Cl³⁷ 2.3%, 584 100%.

20 [3S(1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(2-chlorophenyl)methylthio4-oxopentanoate (224e). 1-Hydroxybenzotriazole (0.23a,

1.71mmol) and ethyl dimethylaminopropyl carbodiimide hydrochloride was added to a stirred solution of the acid 212e (0.295g, 0.853mmol) in THF (5ml). After 5min water (0.5ml) was added followed, after a further 7min. 5 by the addition of a solution of (3S) t-butyl-3allyloxycarbonylamino-5-(2-chloro-phenyl)methylthio-4oxopentanoate (123, 0.478g, 1.02mmol) and (PPh3)2PdCl2 (20mg) in THF (2ml). Tributyltin hydride (0.65ml, 2.33mmol) was added dropwise during 20min. The mixture 10 was kept for 4.5h then diluted with EtOAc, washed with 1M HCl, brine, sat. ac. NaHCO2 and then brine again. The mixture was dried (MgSO₄) and concentrated. The residue was triturated several times with hexane, which was decanted and discarded, then purified by flash 15 chromatography (10-100% EtoAc in CH_2C_{-2}) to afford 0.2g (35%) of a white glassy solid: mp 70-72 °C; $[\alpha]_{D}^{26}$ -82.5 ° (c 0.02, CH₂Cl₂). IR (KBr) 3404, 1726, 1660, 1534, 1524, 1422, 1277, 1254, 1154; ¹H NMR (CDCl₃) δ 7.83-7.78 (2H, m), 7.7, 7.75-7.32, 7.26-7.20 (7H, 3m), 20 7.12 (1H, d, J = 8.2), 7.01 (1H, d, J = 7.3), 5.23-5.08 (2H, m), 5.03-4.94 (1H, m), 4.62 (1H, dt, J = 14.5), 3.78 (2H, m), 3.38-3.29 (1H, m), 3.26 (2H, s), 3.06-2.82 (4H, m), 2.71 (1H, dd, J = 17.2, 4.5), 2.39 (1H, dd, J = 13.2, 6.5), 2.15-1.83, 1.73-1.63 (5H, m), 1.45 25 (9H, s). Anal. Calcd for C33H39ClN4O7S: C, 59.05; H,

[3RS, (1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamido]-5-(2-chlorophenylmethyloxy)-4-30 oxopentanoate (225e), was prepared from acid 212e and (3S) t-butyl N-(allyloxycarbonyl)-3-amino-5-(2-chlorophenylmethyloxy)-4-oxopentanoate (201) using a

5.86; N, 8.35. Found: C, 59.00; H, 5.80; N, 7.92.

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method similar to that used for compound 224e, to afford 40mg (23%) of a glassy solid: $^{1}\text{H NMR}$ (CDCl $_3$) δ 7.83-7.73 (2H, m), 7.67-7.10 (9H, m), 5.23-5.09 (2H, m), 4.59 (1H, m), 4.45-4.22 (2H, m), 3.7-3.19, 3.08-5.72, 2.71-2.47, 2.05-1.85, 1.72-1.61, 1.45-1.26 (20H, 6m).

[38(18,98)] 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamido)-5-(2-chlorophenyl)methylthio-10 4-oxopentanoic acid (226e), was prepared from 224e by an analogous method as that used for compound 217e which afforded 0.22g (81%) of an off-white solid: mp 95-100 °C; [α]_D²³-95.6 ° (c 0.2, CH₂Cl₂). IR (KBr) 3393, 1720, 1658, 1529, 1422, 1279; ¹H NMR (D₆-DMSO) δ 15 8.80 (1H, d, J = 7.5), 7.89 (2H, m), 7.7 (1H, d, J = 7.7), 7.56-7.28 (7H, m), 5.10 (1H, m), 4.87-4.73 (2H, m), 4.39 (1H, m), 3.77 (2H, m), 3.44, 3.35 (2H, +H₂O, 2m), 2.97-2.56, 2.2, 1.92, 1.61 (11H, 4m). Anal. Calcd for C₂₀H₃₁ClM₄O₇S 0.5H₂O: C, 55.02; H, 5.10; N. 8.85.

[3RS, (1S,9S)] 3-Benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(2-chlorophenylmethyloxy)-4oxopentanoic acid (227e), was prepared from 225e by an

25 analogous method as that used for compound 217e. The
product was further purified by flash chromatography
(0-5% MeOH/CH₂Cl₂) to afford 19mg (81.) of a glassy
solid: H NMR (CDCl₃) 8 7.79 (2H, m), 7.66-7.18 (9H,
m), 5.30-5.10 (2H, m), 4.85 (1H, m), 4.65 (2H, m), 4.53

30 (1H, m), 4.28 (2H, m), 3.28, 3.01, 2.72, 2.33, 1.94,
1.60 (1HH, 6m). MS (ES⁻, m/z) 597 (M⁺ - 1, 100°).

20 Found: C, 55.00; H, 5.09; N, 8.71.

229e X =
$$NH \longrightarrow F$$

[3RS,4RS(1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo-

- 5 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamido)-5-fluoro-4(228e). 1-Hydroxybenzotriazole (0.23g, 1.68mmol, followed by ethyldimethylaminopropyl carbodinide hydrochloride (0.21g, 1.09mmol) were added to a stirred
- 10 solution of the acid 212e (0.29g, 0.84mmol) in CH2Cl2 (3ml) at rt. The mixture was kept for 10min them a solution of (3RS,4RS) t-butyl 3-amino-5-fluoro-4-hydroxypentanoate (Revesz, L. et al. Tetrahedror Lett., 52, pp. 9693-9696 (1994); 0.29g, 1.40mmol) in CH2Cl2
- 15 (3ml) was added followed by 4-dimethylaminopyridine

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(10mg). The solution was stirred for 17h, diluted with EtOAc, washed with 1M HCl, brine, sat. aq. NaHCO₃ and brine again, dried (MgSO₄) and concentrated. The residue was purified by flash chromatography (50-100% 5 EtOAc/CH₂Cl₂ and 5% MeOH/EtOAc) to afford 0.25g (56%) of a white glassy solid: IR (KBr) 3343, 1726, 1658, 1536, 1426, 1279, 1257, 1157; ¹H NMR (CDCl₃) 5 7.84-7.79 (2H, m), 7.57-7.40 (3H, m), 7.05-6.92, 6.73 (2H, 2m), 5.17-5.04 (2H, m), 4.56, 4.35-4.21, 4.04 (5H, 3m), 10 3.36, 3.09-2.34, 2.00 (11H, 3m), 1.46 (9H, s). Anal. Calcd for C₂₆H₃₅FN₄O₇ 0.5H₂O: C, 57.45; H, 6.65; N,

[3RS,4RS(1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

15 [1,2-a][1,2]-diazepine-1-carboxamido)-5-fluoro-4oxypentanoate (229e) was prepared from 228c by an
analogous method to that used for compound 216e. After
purification by flash chromatography (30-50%
EtOAc/CH₂Cl₂) the product was obtained as a white

10.31. Found: C, 57.64; H, 6.56; N, 10.15.

20 glassy solid (0.194g, 89%): IR (KBr) 3376, 1728, 1659, 1529, 1424, 1279, 1256, 1156.

[3RS, (1S,9S)] 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamido)-5-fluoro-4-oxopentanoic

- 25 acid (230e), was prepared from 229e by an analogous method to that used for compound 217e to afford 230e as a white glassy solid (100%): mp 105-125 °C; $\{\alpha\}_D^{-23}$ -91.4 ° (c 0.72, CH₃OH). IR (KBr) 3336, 1789, 1737, 1659, 1535, 1426, 1279, 1258, 1186; 1 H NMR (CD₃OD) δ
- 30 7.71-7.68 (2H, m), 7.37-7.23 (3H, m), 5.02, 4.88-4.63,

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4.37-4.0 (6H, 3m), 3.30, 2.97, 2.68-2.60, 2.37-1.54 (11H, 4m). MS(ES, m/z) 475 (M+ - 1, 100%).

(232e)

(231e)

[3S(1S,9S)]-Methyl 9-(benzovlamino)-3-[6,10-dioxo-5 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido]-3-cyanopropanoate (231e). N-Fluorenvlmethvloxy-carbonvl-3-amino-3cyanopropionic acid methyl ester (EP0547699A1, 385mg, 1.1mmol) was treated with 17ml of diethylamine. After 10 1.5h stirring at room temperature the solution was concentrated. The residue was chromatographed on silica gel (3% methanol in CH2Cl2) and gave the free amine as a pale vellow oil. To a solution of this oil and hydroxybenzotriazole (297mg, 2.19mmol) in DMF 15 (5ml), was added at 0 °C ethyldimethylaminopropyl carbodiimide (232mg, 1.21mmol, 1.1 equiv) followed by (15,95) 9-(benzoylamino)-(6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazıno[1,2-a][1,2]diazepine-1carboxylic acid (212e). After stirring at 0 °C for 5 20 min and then at room temperature overnight, the mixture was diluted with CH2Cl2 (50ml) and the resulting solution washed successively with 1M HCl (2 x 30ml). H₂O (30ml), 10% NaHCO₃ (2 x 30ml) and sat. ag. NaCl, dried (MgSO₄) and concentrated. Purification by flash

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chromatography on silica gel (3% methanol in CH_2Cl_2) afforded the compound 231e (404mg, 83%) as a solid: $\left[\alpha\right]_D^{20}$ -121 ° (c 0.14, CH_2Cl_2); ^1H NMR (CDCl₃) 5 7.40-7.83 (5H, m), 7.38 (1H, d), 6.96 (1H, d), 5.27-5.07 5 (2H, m), 4.66-4.50 (1H, m), 3.79 (3H, s), 3.23-2.73 (6H, m), 2.47-2.33 (1H, m), 2.15-1.82 (4H, m); Anal. Calcd for $C_2H_2Sh_5O_6$: C, 58.0; H, 5.53; N, 15.38. Found: C, 57.6; H, 5.6; N, 15.0.

[3S(1S,9S)] 9-(Benzoylamino)-3-[6,10-dioxo-

- 10 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido]-3-cyanopropanoic acid (232e). A solution of methyl ester 231e (400mg, 0.88mmol) in methanol (30ml) and water (30ml) was cooled at 0 °C and treated with diisopropylethylamine.
- 15 The solution was stirred at 0 °C for 10min and then at room temperature overnight. The heterogeneous mixture was concentrated and the solid obtained was chromatographed on silica gel (5% methanol/1% formic acid in CH₂Cl₂) affording the free acid 232e (170mg,
- 20 44%) as a white solid: mp 155 °C (decl; $[\alpha]_D^{20}$ -117 ° (c 0.1, MeOH); IR (KBr) 3343, 3061, 2955, 1733, 1656, 1577, 1533, 1490, 1421, 1342, 1279, 1256, 1222, 1185, 708; 1H NMR (D⁴-MeOH) δ 7.88-7.28 (5H, m), 5.20-5.03 (1H, m), 4.98-4.84 (2H, m), 4.75-4.53 (1H, m), 4.51-
- (H, m), 4.96-4.84 (2H, m), 4.75-4.53 (1H, m), 4.51-25 4.34 (1H, m), 3.45-3.22 (1H, m), 3.14-2.94 (1H, m), 3.14-2.94 (1H, m), 2.88-2.61 (2H, m), 2.53-1.50 (8H, m); Anal. Calcd for C₂₁H₂₂N₅O₆. 1.5H₂O: C,53.84; H, 5.59; N, 14.95; C, 25.61. Found: C, 54.3; H, 5.4; N, 14.3.

[45, (15,95)] t-Butyl 4-[9-(benzoylamino)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido]-5-oxopentanoate semicarbazone (233e). A sclution of (15,95) 6,10-5 dioxo-1,2,3,4,7,8,9,10-octahydro-9-(benzoylamino:-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxylic acid (212e) (345mg, 1.0mmol), (45) t-butyl N-

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(allyloxycarbonyl)-4-amino-5-oxopentanoate semicarbazone (208a) (361mg, 1.1mmol, 1.1 equiv) and $(Ph_3P)_2PdCl_2$ (20mg) in CH_2Cl_2 (5ml), was treated dropwise with n-Bu₃SnH (0.621ml, 2.3mmol, 2.1 equiv).

- 5 The resulting orange brown solution was stirred at 25 °C for 10min and then 1-hydroxybenzotriazole (297mg, 2.2mmcl, 2 equiv) was added. The mixture was cooled to 0 °C and ethyldimethylaminopropyl carbodiimide (253mg, 1.3mmol, 1.2 equiv) added. After stirring at 0 °C for
- 10 10min and then at room temperature overnight, the mixture was diluted with EtOAc (50ml) and the resulting solution washed successively with 1M HCl (3 x 25ml), 10% NaHCO₃ (3 x 25ml) and sat. aq. NaCl, dried (MgSO₄) and concentrated. Flash chromatography on silica gel
- 15 (2-10% methanol in $\mathrm{CH_2Cl_2}$) afforded compound 233e (280mg, 49%) as a tan solid: $\left[\alpha\right]_{D}^{20}$ -95 (c 0.09, MeOH); IR (KBr) 3477, 3333, 2968, 2932, 1633, 1580, 1535, 1423, 1378, 1335, 1259, 1156, 1085, 709; $^1\mathrm{H}$ NMR (CDCl₃) δ 9.32 (1H, s), 7.83-7.39 (6H, m), 7.11-7.09 (1H, m).
- 20 6.30-5.30 (2H, brs), 5.17-5.05 (2H, m), 4.62-4.38 (2H, m), 3.30-3.15 (1H, m), 3.13-2.65 (2H, m), 2.46-2.19 (3H, m), 2.15-1.54 (8H, m), 1.42 (9H, s).

[4R, (1S,9S)] t-Butyl 4-[9-(benzoylamino)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

- 25 [1,2-a][1,2]diazepine-1-carboxamido]-5-oxopentanoate
 semicarbazone (236e), was prepared by an analogous
 method to that used for 233e using (4R) t-butyl N allyloxycarbonyl-4-amino-5-oxo-pentanoate semicarbazone
 (208b, 435mg, 1.33mmol). The product was obtained as a
 30 foam (542mg, 71½): [α]_D²⁰ -99 ° (c 0.19, CHCl₃); IR
 (KBr) 3473, 3331, 3065, 2932, 2872, 1660, 1580, 1533,
 - 1488, 1423, 1370, 1337, 1278, 1254, 1223, 1155, 1080,

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1024, 983, 925, 877, 846, 801, 770, 705; ¹H NMR (CDCl₂) 5 9.42 (1H, s), 7.81 (2H, d), 7.51-7.40 (4H, m), 7.06 (1H, d), 6.50-5.50 (2H, broad s), 5.25-5.00 (2H, m), 4.60-4.45 (2H, m), 3.15-2.85 (2H, m), 2.75-2.35 (1H, m), 2.30-1.23 (11H, m), 1.42 (9H, s).

[4s, (1s,9s)] t-Butyl 4-[9-(benzoylamino)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido]-5-oxopentanoate (234e). A solution of semicarbazone 233e (390mg.

- 10 0.68mmol) in methanol (10ml) was cooled at 0 °C and then treated with a 38% aq. solution of formaldehyde (2ml) and 1M HCl (2ml). The reaction mixture was then stirred overnight at room temperature. The solution was concentrated to remove the methanol. The aq.
- 15 solution was extracted with EtOAc (30ml). The organic solution was successively washed with 10% NaHCO₃ (30ml) and sat. aq. NaCl (30ml), dried (MgSO₄) and concentrated. Purification by flash chromatography on silica gel (2-5% methanol in CH,CL₂) afforded 234e
- - [4R, (1S,9S)] t-Butyl 4-[9-(benzoylamino)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido]-5-oxopentanoate (237e), was prepared from 236e by an analogous method
- 30 (237e), was prepared from 236e by an analogous method to 234e to afford a white foam (390mg, 85%): $[\alpha]_{\rm p}^{20}$

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-113 ° (c 0.242, CHCl₃); IR (KBr) 3352, 3065, 2974, 1729, 1657, 1536, 1489, 1454, 1423, 1369, 1338, 1278, 1255, 1223, 1156, 1078, 1026, 981, 846, 709.

[4S, (1S,9S)] 4-[9-(Benzoylamino)-6,10-dioxo-

- 5 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-
 - [1,2-a][1,2]diazepine-1-carboxamido]-5-oxopentanoic acid (235e). A solution of t-butyl ester 234e (179 $_{
 m mg}$, 0.35 $_{
 m mmol}$) in dry CH $_{
 m 2}$ Cl $_{
 m 2}$ (3 $_{
 m ml}$) was cooled to 0 °C and
- treated with trifluoroacetic acid (2ml). The resulting 10 solution was stirred at 0 °C for 30min and then at room temperature for 2h. The solution was concentrated, the residue taken up in dry $\mathrm{CH_2Cl_2}$ (5ml) and the mixture again concentrated. This process was repeated once
- again with more ${\rm CH_2CI_2}$ (5ml). The residue obtained was 15 crystallized in diethyl ether. The precipitate was collected and purified on silica gel column (5%)
- collected and purified on silica gel column (5% methanol in CH_2Cl_2) which afforded compound **235e** as a white solid (111mg, 70%): mp 142 °C (dec); $\{\alpha\}_D^{20} = 85.5$ (c 0.062, MeOH); IR (KBr) 3409, 3075, 2952, 1651, 1541,
- 20 1424, 1280, 1198, 1136, 717; ¹H NMR (D₆-DMSO) δ 9.40 (1H, s), 8.62 (2H, m), 7.96-7.38 (5H, m), 5.19-5.02 (1H, m), 4.98-4.79 (1H, m), 4.48-4.19 (1H, m), 3.51-3.11 (2H, m), 3.04-2.90 (2H, m), 2.38-1.46 (10H, m).

[4R, (1S,9S)] 4-[9-(Benzoylamino)-6,10-dioxo-

25 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-5-oxopentanoic
 acid (238e), was prepared from 237e by an analogous
 route to 235e which afforded a beige foam (190mg, 60%):
 [α]_D²⁰ -78 (c 0.145, MeOH); lR (KBr) 3400, 3070, 2955.
30 2925, 2855, 1653, 1576, 1541, 1490, 1445, 1427, 1342.

1280, 1258, 1205, 1189, 1137, 1075, 1023, 983, 930,

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878, 843, 801, 777, 722; ¹H NMR (D₆-DMSO) 8 9.40 (1H, s), 8.72-8.60 (2H, m), 7.89 (2H, d), 7.56-7.44 (3H, m), 5.17 (1H, m), 4.90-4.83 (1H, m), 4.46-4.36 (1H, m), 4.20-4.15 (1H, m), 3.40-3.30 (1H, m), 2.98-2.90 (2H, m), 2.50-1.60 (10H, m).

(15,95) t-Butyl 9-benzoylamino-octahydro-10-oxo-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxylate (243),

10 was prepared from (15,95) t-butyl 9-amino-octahydro-10-oxo-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxylate (Attwood, et al. J. Chem. Soc., Perkin 1, pp. 1011-19 (1986)), by the method described for 211e, to afford 2.03g (86%) of a colourless foam: [α]₀²⁵ -15.9 ° (c
15 0.5, CH₂Cl₂); IR (KBr) 3400, 2976, 2937, 174C, 1644, 1537, 1448, 1425, 1367, 1154; ¹H NMR (CDCl₃) δ 7.88-7.82 (2H, m), 7.60-7.38 (4H, m), 5.46 (1H, n), 4.98 (1H, m), 3.45 (1H, m), 3.22-2.96 (2H, m), 2.64 (1H, m), 2.43-2.27 (2H, m), 1.95 (2H, m), 1.82-1.36 (4H, m), 2.61 (1H, m), 2.91 (1H, m), 3.75 (1H, m), 2.92 (1.50 (9H, s); Anal. Calcd for C₂₁H₂₉N₃O₄, 0.25H₂O; C,

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64.35; H, 7.59; N, 10.72. Found: C, 64.57; H, 7.43; N, 10.62. MS (ES +, m/z) 388 (100%, M^* + 1).

(1S,9S) 9-Benzoylamino-octahydro-10-oxo-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxylic acid
5 (244), was prepared from (1S,9S) t-butyl 9-benzoylamino-octahydro-10-oxo-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxylate (243), by the method described for 212e, to afford 1.52g (89%) of a white powder: mp. 166-169 °C (dec); [α]_D²⁵ -56.4 ° (c
10 0.5, CH₉OH); IR (KBI; 3361, 2963, 2851, 1737, 1663, 1620, 1534, 1195, 1179; ¹H NMR (D₆-DMSO) δ 12.93 (1H, brs), 8.44 (1H, d, J = 8.4), 7.93 (2H, m), 7.54 (3H, m), 5.46 (1H, m), 4.87 (1H, m), 3.12 (2H, m), 2.64 (1H, m), 2.27 (1H, m), 1.98-1.68 (7H, m), 1.40
15 (1H, m); Anal. Calcd for C₁₇H₂₁N₃O₄. 0.25H₂O: C, 60.79; H, 6.45; N, 12.51. Found: C, 61.07; H, 6.35; N.

[3s,2Rs(1s,9s)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-9-benzovlamino-octahydro-10-oxo-6H-

12.55. MS (ES+, m/z) 332 (58%, M+ + 1), 211 (100).

- 30 521 (100\, M + 1).

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[3S(1S,9S)] 3-(9-Benzoylamino-octahydro-10-oxo-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide-4-oxobutanoic acid (246), was prepared from [3S, 2RS (1S,9S)]N-(2-benzyloxy-5-oxotetrahydrofuran-3-yl)-9-

- 5 benzoylamino-octahydro-10-oxo-6H pyridazino[1,2-a][1,2]diazepine-1-carboxamide (245), by
 the method described for 214e, to afford 396mg (84%) of
 a white powder: mp. 110-115 °C; (α)_D²⁶ -126.3 ° (c 0.2,
 CH₃OH); IR (KBr) 3345, 2943, 1787, 1730, 1635, 1578,
- 10 1528, 1488, 1450, 1429; ^{1}H NMR (CD₃OD) δ 7.88 (2H, m), 7.48 (3H, m), 5.55 (1H, m), 4.91 (1H, m), 4.56 (1H, m), 4.29 (1H, m), 3,41-3.05 (3H, m), 2.76-2.41 (3H, m), 2.8-2.01 (3H, m), 1.86-1.65 (4H, m), 1.36 (1H, m); Anal. Calcd for C₂₁H₂₆N₄O₆. 1.25H₂O: C, 55.68; H, 6.34; N, 1.37, Calcal Calca
- 15 N, 12.37. Found: C, 55.68; H, 6.14; N, 12.16. MS (ES -, m/z) 429 (100%, M^{+} 1).

[(3S(2R, 5S)]-2,6-Di-tert-buty1-4-methoxypheny1-3-[5-(2,5-dihydro-3,6-dimethoxy-2-(1-

5 methylethyl)pyrazinyl)]butanoate (247). n-Butyllithium (1.6M in hexane) (22.3ml, 35.7mmol) was added dropwise over 20min to a solution of (2R)-(-)-2,5-dihydro-3,6-dimethoxy-2-(1-methylethyl)pyrazine (5.8ml, 6.0g, 32.4mmol) in THF (250ml) cooled to -75 °C at a rate such that the temperature was maintained below -72 °C. The reaction mixture was stirred for 1h at -75 °C and a solution of 2,6-di-t-butyl-4-methoxyphenyl-2-butenoate (Suzuck et al. Liebigs Ann. Chem. pp. 51-61 (1992))

(9.9g, 32.5mmol) in THF (60ml) was added over 30 minutes maintaining the temperature below -72 °C during the addition. The reaction mixture was kept at -75 °C for 1.5h and a solution of glacial acetic acid (6ml) in 5 THF (25ml) was added at -75 °C and the solution warmed to room temperature. The solution was poured onto 10% NH₄Cl (300ml) and extracted with diethyl ether (3 x 250ml). The combined organic phases were washed with brine (2 x 200ml), dried over Na2SO4 and evaporated to 10 dryness under reduced pressure. The residual oil was purified by flash chromatography on silica gel (20% heptane in CH2Cl2) which afforded the title compound as a light yellow oil (13.5g, 85%): $[\alpha]_{\rm p}^{20}$ -64 ° (c 0.22, MeOH); IR (KBr) 2962, 2873, 2840, 1757, 1697, 1593, 15 1460, 1433, 1366, 1306, 1269, 1236, 1187, 1157, 1126, 1063, 1038, 1011, 970, 924, 892, 867, 846, 831, 797, 773, 754; ¹H NMR (CDCl₃) δ 6.85 (2H, s), 4.21 (1H, t, J = 3.5), 3.98 (1H, t, J = 3.5), 3.79 (3H, s), 3.71 (3H, s), 3.69 (3H, s), 3.15 (1H, dd, J 17.8, 7.9),

20 2.86-2.81 (1H, m), 2.58 (1H, dd, J = 17.8, 5.9), 2.28-2.19 (1H, m), 1.33 (18H, s), 1.02 (3H, d, J = 6.8), 0.70 (6H, dd, J = 13, 6.8).

(25,38)-5-[2,6-Di-t-butyl-4-methoxyphenyl]1-methyl-3-methylglutamate (248). A solution of [35(2R, 55)]-2,6-25 di-t-butyl-4-methoxyphenyl-3-[5-(2,5-dihydro-3,6-dimethoxy-2-(1-methylethyl)pyrazinyl)]butanoate (247) (22.4g, 45.8mmol) in acetonitrile (300ml) and 6.25N HCl (366ml, 2 equiv) was stirred at room temperature under nitrogen atmosphere for 4 days. The acetonitrile was evaporated under reduced pressure and diethylether (250ml) was added to the aq. phase. The pH of the aq. phase was adjusted to pH8-9 with concentrated ammonia

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solution (32%) and the phases separated. The aq. phase was extracted with diethylether (2 x 250ml). The combined organic phases were dried over Na_2SO_4 and evaporated to dryness under reduced pressure. The

- 5 residual oil was purified by flash chromatography on silica gel (2% methanol in CH₂Cl₂) which afforded the required product as a light yellow oil (8.2g, 45%): [a]_D²⁰ +20° (c 0.26, MeOH); IR(KBr) 3394, 3332, 3000, 2962, 2915, 2877, 2838, 1738, 1697, 1593, 1453, 1430,
- 10 1419, 1398, 1367, 1304, 1273, 1251, 1221, 1203, 1183, 1126, 1063, 1025, 996, 932, 891, 866, 847, 800, 772, 745; ^{1}H NMR (CDCl₃) δ 6.85 (2H, s), 3.79 (3H, s), 3.74 (3H, s), 3.72-3.69 (1H, m), 3.05-2.85 (1H, m), 2.67-2.50 (2H, m), 1.32 (18H, s), 0.93 (3H, d, J = 7); Anal.
- 15 Calcd for $C_{22}H_{35}NO_5$: C, 67.15; H, 8.96; N, 3.56. Found: C, 67.20; H, 9.20; N, 3.70.

(2s,3s)-5-[2,6-Di-t-butyl-4-methoxyphenyl]3-methylglutamate (249). A solution of (2s,3s)-5-[2,6-di-t-butyl-4-methoxyphenyl]3-methylglutamate (248)

- 20 (8.0g, 20.3mmol) in 5N HCl (200ml) was heated at reflux for 2h. The reaction mixture was evaporated to dryness under reduced pressure. The residue was dissolved in cyclohexane (x4) and evaporated to dryness (x4) which afforded a white solid (7.9g, 93%): mp 230 °C; [0]p²⁰
 25 +22 ° (c 0.27, MeOH); IR (KBr) 3423, 2964, 1755, 1593,
- 25 +22 ° (c 0.27, MeOH); IR (KBr) 3423, 2964, 1755, 1593, 1514, 1456, 1421, 1371, 1303, 1259, 1201, 1179, 1138, 1106, 1060, 966, 926, 861, 790, 710; ¹H NMR (MeOD) 8 6.76 (2H, s), 4.02 (1H, d, *J* = 3.7), 3.67 (3H, s), 3.05-2.85 (1H, m), 2.80-2.55 (2H, m), 1.22 (18H, s),
- 30 1.09 (3H, d, J = 6.3); ¹³C NMR (MeOD) δ 174.5, 171.4, 158.6, 145.2, 143.1, 113.2, 58.3, 56.3, 39.8, 36.9,

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32.5, 16.6; Anal. Calcd for $C_{21}H_{34}ClNO_5$: C, 60.64; H, 8.24; N, 3.37. Found: C, 60.80; H, 8.40; N, 3.40.

(2S,3S)-5-[2,6-Di-t-butyl-4-methoxyphenyl]3-methyl-2-phthalimido-1,5-pentanedioate (250),

- 5 Diisopropylethylamine (4.1ml, 3.04g, 23.5mmol, 1.25 equiv) and phthalic anhydride (3.5g, 23.6mmol, 1.25 equiv) were added to a solution of (25,35)-5-[2,6-di-t-butyl-4-methoxyphenyl]3-methylglutamate (249) (7.8g, 18.6mmol) in toluene (300ml). and the resulting mixture
- 10 was heated at reflux for 3 hours. After cooling to room temperature, the reaction mixture was evaporated to dryness and the resulting oil purified by flash chromatography on silica gel (2% methanol in CH2Cl2) which afforded the required product as a white foam
- 20 3.30-3.05 (2H, m), 2.85-2.65 (1H, m), 1.30 (18H, s), 1.13 (3H, d).

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10 (257) (256)

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- 1-(2,6-di-t-Butyl-4-methoxy)-phenyl-5-(1-benzyloxycarbonyl-3-t-butoxycarbonyl-hexahydro-pyridazin-2-yl)-3-methyl-4-phthalimidopentan-1,5-dioate (251). A solution of the amino acid (250) (1.2q,
- 5 2.35mmol) in dry diethylether (10ml) was treated with phosphorus pentachloride (0.52g, 2.5mmol) at room temperature for 2h. The mixture was concentrated and treated several times with toluene and again evaporated to dryness. The resulting acid chloride was dissolved
- 10 in dry THF (5ml) and CH₂Cl₂ (5ml) and cooled to 0 °C. t-Butyl-1-(benzyloxycarbonyl)-hexahydro-3-pyridazinecarboxylate (0.753g, 2.35mmol, 1 equiv) and Nethylmorpholine (3ml) were added to the solution. The reaction mixture was stirred for 30min at 0 °C and then
- 15 overnight at room temperature. The mixture was evaporated and the resulting residue taken up with $\mathrm{CH_2Cl_2}$ (30ml). The solution was washed with 1M HCl, water, 10% NaHCO₃, dried (MgSO₄) and evaporated. The resulting white foam was purified on silica gel (0-2% or 10-2% o
- 20 methanol in CH₂Cl₂) which afforded the required
 compound 251 as a pale yellow glassy solid (740mg,
 39%): [α]_D²⁰ -22 (c 0.42, MeOH); IR (KBr) 3441, 2966,
 1725, 1693, 1386, 1255, 1221, 1186, 1154, 1123, 1063,
 724; ¹H NMR (CDCl₃) δ 7.94-7.89 (4H, m), 7.56-7.28 (5H,
- 25 m), 6.84 (2H, 2s), 5.29-5.20 (2H, AB), 4.91-4.81 (1H, m), 4.05-3.88 (1H, m), 3.78 (3H, s), 3.75-3.80 (1H, m), 3.28-2.95 (2H, m), 2.23-1.51 (6H, m), 1.45 (9H, s), 1.31 (9H, s), 1.20 (9H, s), 1.27 (3H, d).
 - (1S, 8S, 9S) t-Butyl 6,10-dioxo-8-methyl-
- 30 1,2,3,4,7,8,9,10-octahydro-9-phthalimido-6H-pyridazino[1,2-a][1,2]diazepin-1-carboxylate (254). A solution of the protected acid (251) (715mg, 0.893mmol)

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in acetonitrile was treated with Cerium (IV) ammonium nitrate (1.8g, 3.3mmol, 3.7 equiv) in water (3ml) for 4h at room temperature. Mannitol (600mg, 3.3mmol, 3.7 equiv) was added and the mixture was stirred for 1h. 5 Diethylether (50ml) and water (30ml) were added to the mixture. After decantation, the ag. phase was extracted with diethylether (4 x 50ml). The combined organic phase was washed with water, dried (MgSO₄) and concentrated. Chromatography on silica gel (10% 10 methanol in CH2Cl2) afforded 5-(1-benzyloxycarbonyl-3t-butoxycarbonyl-hexahydropyridazin-2-yl)carbonyl-3methyl-4-phthalimidopentanoic acid (252) (360mg, 64%): $\lceil \alpha \rceil_n^{20}$ -49.2 c 0.118, MeOH). This product was used without further purification (360mg, 0.609mmol), and 15 was hydrogenated in methanol (30ml) using 10% Pd/carbon (36mg) for 3h. The reaction mixture was filtered and the resulting solution concentrated to afford the amine (253) as a foam (270mg, 96%) $\left[\alpha\right]_{D}^{20}$ -56.1 (c 0.18 MeOH). The amine (253) was dissolved in dry THF (10ml) and 20 phosphorous pentachloride (305mg, 1.47mmcl, 2.5 equiv) was added. The mixture was then cooled to -5 °C and Nethylmorpholine was added under nitrogen. The reaction mixture was stirred overnight at room temperature. The mixture was concentrated and the residue taken up with 25 CH₂Cl₂ (20ml), cold H₂O (20ml), 1M HCl (20ml). After decantation, the ag. phase was reextracted with CHoClo (2 x 20ml). The combined organic phase was washed with 10% NaHCO3 and water, dried (MgSO4) and concentrated. The resulting oil was purified on silica gel (1% 30 methanol in CH2Cl2) affording the bicyclic compound (254) as a solid (65mg, 25%): $[\alpha]_{\rm p}^{20}$ -77 (c 0.20%, MeOH): IR (KBr) 3471, 3434, 2975, 2928, 1767, 1723, 1443, 1389, 1284, 1243, 1151, 1112, 720; ¹H NMR (CDC1₂₁

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(1S, 8S, 9S) t-Butyl-9-benzoylamino-6,10-dioxo-8-methyl-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-

[1,2-a][1,2]diazepine-1-carboxylate (255). A solution of the bicyclic compound (254) (70mg, 0.16mmol) in 10 ethanol was treated with hydrazine hydrate (0.02ml. 4mmol, 2.5 equiv). After 5h stirring at room temperature, the mixture was concentrated and the resulting residue taken up in toluene and reevaporated. The residue was treated with 2M acetic acid (2ml) for 15 16h. The resulting precipitate was filtered and washed with 2M acetic acid (10ml). The filtrate was basified with solid NaHCO3 and then extracted with EtOAc. The organic solution was washed with water, dried (MgSO₄) and concentrated. Purification by flash chromatography 20 on silica gel (2% methanol in CH2Cl2) afforded the free amine as a foam (50mg, 100%). The amine (50mg, 0.16mmol) was dissolved in dioxane (1ml) and water (0.25ml) and treated with NaHCO3 (0.034g, 0.04mmol) followed by benzoylchloride (0.047ml, 0.40mmol, 2.8 25 equiv). The mixture was stirred overnight at room temperature, then diluted with EtOAc (15ml). The organic solution was washed with 10 - NaHCO3 and sat. ag. NaCl, dried (MgSO4) and concentrated. Purification by flash chrcmatography on silica gel (2: methanol 17 30 CH₂Cl₂) afforded the benzamide 255 as a foam (67mg, 100%): ¹H NMR (CDCl₃) 8 7.89-7.39 (5H, m), 6.79 (1H, d), 5.32-5.20 (1H, m), 4.98-4.82 (1H, m), 4.75-4.64

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(1H, m), 3.84-3.65 (1H, m), 3.09-2.89 (1H, m), 2.45-2.18 (2H, m), 2.00-1.61 (4H, m), 1.48 (9H, s), 1.28 (3H, d).

[3S(1S, 8S, 9S)] 3-(9-benzovlamino-6,10-dioxo-8-methyl-5 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (257). A solution of t-butyl ester 255 (67mg, 0.16mmol) in CH2Cl2 (1ml) was treated at 0 °C with trifluoroacetic acid (lml). The resulting solution was 10 stirred at 0 °C for 15min and then at room temperature for in. The solution was concentrated, the residue taken up in dry CH2Cl2 (2 x 2ml) and the mixture again concentrated (x2). The residue was crystallized from diethylether. Filtration of the precipitate afforded 15 the free acid of 255 as a grey solid (40mg, 70%). A solution of acid (40mg, 0.11mmol), N-allyloxycarbonyl-4-amino-5-benzyloxy-2-oxotetrahydrofuran (Chapman, Biocrg. & Med. Chem. Lett., 2, pp. 615-18 (1992); 39mg, 9.13mmol, 1.2equiv) and (PhaP)aPdCla (3mg) in a mixture 20 of dry CH₂Cl₂ (1ml) and dry DMF (0.2ml) was treated dropwise with n-Bu3SnH (0.089ml, 0.33mmol, 3 equiv). The resulting solution was stirred at 25 °C for 10min and then 1-hydroxybenzotriazole (36mg, 0.266mmol, 2.4 equiv) was added. The mixture was cooled to 0 °C and 25 ethyldimethylaminopropyl carbodiimide (31mg, 6.16mmol, 1.5equiv) was added. After stirring at 0 °C for 10min and then at room temperature overnight, the mixture was diluted with EtOAc (20ml) and the resulting solution washed successively with 1M HCl (2 x 5ml), 10% NaHCO2 30 (2 x 5ml) and sat. aq. NaCl (5ml), dried (MgSCz) and concentrated. Flash chromatography on silica gel (2 methanol in CH2Cl2) afforded a mixture of

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diastereoisomers (256) as a grey solid (50mg, 82%). This product (256) was used without further purification (50mg, 0.091mmol) and was hydrogenated in methanol (5ml) using 10% Pd/carbon (30mg) for 24h. The reaction mixture was filtered and the resulting solution concentrated. Flash chromatography on silica gel (2-20% methanol in CH₂Cl₂) afforded compound 257 (9mg, 21%) as a white solid: ¹H NMR (D⁴-MeOH) & 7.88-7.29 (5H, m), 5.18-4.99 (1H, m), 4.59-4.35 (3H, m), 4.26-4.11 (1H, m), 3.65-3.41 (2H, m), 3.18-2.91 (1H, m), 2.62-1.47 (8H, m), 1.29-1.00 (3H, 2d) (mixture of

acetal and hemiacetal). MS (ES -) 457.

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5

Benzyl 3-(N'-benzoylhydrazino)propanoate (259).

Benzylacrylate (1.13ml, 7.34mmol) was added to a

10 stirred suspension of benzoylhydrazine (285) (1.0g,
7.34mmol) in isopropanol (28ml). The mixture was

refluxed for 20h, cooled to room temperature then concentrated. The residue was purified by flash chromatography (20% EtoAc in CH₂Cl₂) to afford **259** (1.098g, 50%) as an oil which crystallized on standing: 5 mp 65 °C; IR (KBr) 3283, 1723, 1644, 1316, 1201, 1156; ¹H NMR (CDCl₃) & 8.32-8.18 (1H, m), 7.81-7.70 (2H, m), 7.57-7.23 (8H, m), 5.36-4.92 (1H, brm), 5.11 (2H, s), 3.26 (2H, t, J = 6.5), 2.59 (2H, t, J = 6.5); ¹³C NMR (CDCl₃) & 172.12, 167.27, 135.65, 132.54, 131.66, 10 128.45, 128.10, 128.06, 126.84, 66.31, 47.33, 33.31; Anal. Calcd for Cl₁H₁₈N₂O₃: C, 68.44; R, 6.08; N, 9.39. Found: C, 68.42; H, 6.10; N, 9.36. MS (ES +) 321 (M + Found: C, 68.42; H, 6.10; N, 9.36; MS (ES +) 321 (M + Found: C, 68.42; H, 6.10; N, 9.36

(3S)-1-Benzyl 3-t-butyl 2-(N'-benzoyl-N-(2-

Na, 38%), 299 $(M^+ + 1, 100)$.

- benzyloxycarbonylethyl)hydrazinocarbonyl)hexahydropyridazine-1,3-dicarboxylate (260). A solution of (35)-1-benzyl 3-t-butyl hexahydropyridazine-1,3dicarboxylate (Hassall et al. J. Chem. Soc. Perkin 1, pp. 1451-1454 (1979)) (925.3mg, 2.89mmol) and
- 20 diisopropylethylamine (0.70ml, 4.0mmol) in a 1.93M toluene solution of phosgene (17.96ml, 34.7mmol) was stirred at room temperature for 45min, then concentrated to leave a yellow solid. To this solid was added toluene (18ml), hydrazide (259) (861.6mg,
- 25 2.89mmol) and diisopropylethylamine (0.70ml, 4.6mmol). The mixture was stirred at room temperature for 2.75n, then concentrated. The resulting residue was taken up in EtOAc, washed twice with 1M HCl, brine, then dried (MgSO₄), filtered and concentrated to afford 2.15g cf
- 30 crude material. Flash chromatography (40° EtoAc in hexane) afforded 1.65g (89%) of the title compound as a white foam: mp 40 °C; [α]p24 -55.78 ° (c 0.40, CH_CCl₂);

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IR (KBr) 3436, 2930, 1733, 1689, 1455, 1412' 1367, 1258, 1156, 697; 1 H NMR (CDCl₃) & 8.54-8.23 (0.5H, m), 7.97-7.09 (15.5H), 5.16-4.80 (4H, m), 4.66-4.32 (1H, m), 4.24-3.55 (3.3H, m), 3.50-3.26 (0.4H, m), 3.19-2.49 (2.3H, m), 2.11-1.43 (6H, m), 1.32-1.05 (7H, m); Anal. Calcd for $C_{35}H_{40}N_{4}O_{6}$ ·0.5H₂O: C, 64.31; H, 6.32; N, 8.57. Found: C, 64.18; H, 6.27; N, 8.56. MS (ES+) 662 (M + Na, 848), 645 (M' + 1, 100), 384 (77).

(6S) -3-(N'benzoyl-N-(6-t-butoxycarbonylhexa-

- hydropyridazine-1-carbonyl)hydrazino)propanoic acid (261). A solution of 260 (1.59g, 2.47mmol) in MeOH (142ml) was treated with 10% Palladium on carbon (230.0mg) and stirred under an atmosphere of $\rm H_2$ for 1.5h. The mixture was filtered and the solvent
- 15 evaporated to afford 1.04g (100%) of a white foam. This was used in the next step without further purification: mp <40 °C; $[\alpha]_D^{26} + 1.6$ ° (c 0.26, CH₂Cl₂); IR (KBr) 3422, 2977, 2986, 1728, 1677, 1486, 1445, 1396, 1369, 1309, 1228, 1155, 916, 716; $^1_{\rm H}$ NMR
- 20 (CDCl₃) δ 10.0-9.7 (1H, brm), 7.86 (2H, d, J = 7.5), 7.62-7.38 (3H, m), 7.3-5.6 (2H, brm), 4.57 (1H, brd, J = 4.0), 4.05-3.77 (2H, m), 3.00-2.82 (1H, m), 2.80-2.43 (3H, m), 2.20-2.03 (1H, m), 2.00-1.47 (1H, m), 1.62-1.14 (11H, m); 13 C NMR (CDCl₃) δ 175.00, 171.17, 167.62,
- 25 160.68, 132.39, 131.77, 128.67, 127.38, 82.27, 54.38, 48.04, 46.35, 33.62, 28.02, 25.68, 21.61. MS (ES -) 443 (M + Na, 68%), 421 (M⁺ + 1), 100), 365 (50), 131 (61).
- (4S) t-Butyl 7-benzamido-6,10-dioxo-1,2,3,4,7,8,9,10-30 octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4carboxylate (262). To a solution of amino acid 261

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(1.012g, 2.41mmol) in dry THF (26ml) at 0 °C was added N-ethylmorpholine (597µl, 4.69mmol), followed by PCl $_5$ (651.3mg, 3.12mmol). The reaction was stirred at 0 °C for 2h, then allowed to warm to rt and stirred for a

- 5 further 15.5h. The mixture was concentrated and the resulting residue taken up in EtOAc, washed twice with 1M HCl, sat. NaHCO₃, brine, then dried (MgSO₄), filtered and concentrated. Flash chromatography (20% EtOAc in CH₂Cl₂) gave 727.3mg (75%) of the title
- 10 compound as a white foam: $(\alpha)_0^{26} + 51.0^{\circ}$ (c 0.20, CH₂Cl₂); IR (KBr) 3436, 2979, 1733, 1670, 1483, 1437, 1420, 1299, 1243, 1156; 1 H NMR (CDCl₃) δ 8.70 (1H, s), 7.78 (2H, d, J = 7.0), 7.57-7.32 (3H, m), 5.08 (1H, dd, J = 2.5, 5.5), 4.59-4.43 (1H, m), 4.08-3.69 (3H, m),
- 15 3.07-2.84 (1H, m), 2.57-2.35 (1H, m), 2.34-2.14 (1H, m), 2.07-1.43 (3H, m), 1.48 (9H, s); ¹³C NMR (CDCl₃) 8 172.41, 169.04, 166.35, 158.35, 132.24, 132.03, 128.61, 127.31, 82.77, 55.41, 54.07, 41.57, 32.21, 28.04, 24.97, 20.37; Anal. Calcd for ConHockley C. C, 59.69; H,
- 20 6.51; N, 13.92. Found: C, 59.53; H, 6.53; N, 13.84.

 MS (ES +) 425 (M + Na, 71%), 403 (M+ + 1, 100), 145

 (41).

(4S)-7-Benzamido-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic Acid

- 25 (263). A solution of ester 262 (720.0mg, 1.80mmol) in a 1:1 mixture of $\mathrm{CH_2Cl_2}$ and TFA (150ml) was stirred for 1.3h under a dry atmosphere. The solution was then reduced in vacuo, taken up in $\mathrm{Et_2O}$ and reduced again. This process was repeated six times to afford the crude
- 30 product as an off-white solid. The product was purified by flash chromatography (5% MeOH in $\rm CH_2Cl_2$) to afford 520.0mg (83%) of the title compound as a white

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foam: $[\alpha]_D^{25} + 59.5^{\circ}$ (c 1.82, CH₂Cl₂); IR (KBr) 3435, 3266, 2956, 1732, 1664, 1524, 1486, 1440, 1302; 1 H NMR (CDCl₃) δ 9.13 (1H, s), 7.77 (2H, d, J = 7.5), 7.57-7.32 (3H, m), 5.27-5.16 (1H, m), 4.62-4.43 (1H, m), 4.09-2.70 (3H, m), 3.14-2.89 (1H, m), 2.59-2.43 (1H, m), 2.38-2.20 (1H, m), 2.14-1.89 (1H, m), 1.82-1.59 (2H, m); 13 C NMR (CDCl₃) δ 173.65, 172.28, 166.44, 158.42, 132.44, 131.31, 128.61, 127.39, 54.83, 54.01, 42.11, 31.79, 24.42, 20.29; MS (ES -) 345 (M - H⁺, 10 1008), 161 (45).

[2RS,3S(4S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxamide (264). To a solution of acid 263 (300.0mg, 0.87mmol) and

- The reaction was stirred for 5min, then 1hydroxybenzotriazole (234.lmg, 1.73mmol) was added and
 the mixture was cooled to 0 °C before addition of 1-(3dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride
 25 (204.5mg, 1.04mmol). The mixture was allowed to warm
- to rt and stirred for 16.5h. The mixture was allowed to warm with EtOAc, washed with 1M NaHSO₄ twice with sat. NaHCO₃, then H₂O and brine. The organic layer was dried (MgSO₄), filtered and concentrated. The residue
- 30 was purified by flash chromatography (5: MeOH in CH₂Cl₂) to afford 358.3mg (77%) of the title compound as a white solid: IR (KBr) 3435, 1791, 1665, 1526,

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1421, 1285; ^{1}H NMR (CDCl₃) 3 8.76 and 8.49 (1H, 2 x s), 7.92-7.73 (2H, m), 7.62-7.24 (8.5H, m), 6.86 (0.5H, d, J=8.0), 5.53 and 5.33 (1H, d, J=5.5, s), 4.95-4.34 (5H, m), 4.04-3.54 (3H, m), 3.03-2.64 (2H, m), 2.49-5 2.14 (2H, m), 2.11-1.46 (4H, m); MS (ES +) 558 (M + Na, 100%), 536 (M + 1, 78), 404 (58).

[35(45)]3-(7-Benzamido-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido)-4-oxobutanoic acid (265). A mixture of

- 10 **264** (350.0mg, 0.65mmol), 10% palladium on carbon (350mg) and methanol (36ml) was stirred under an atmosphere of $\rm H_2$ for 6.5h. The mixture was filtered and the solvent evaporated. Et $_2$ O was added and the solvent removed again. This process was repeated four
- 15 times to reveal 283mg (97%) of the title compound, as a white crystalline solid: mp decarboxylates above 140 °C; $[\alpha]_{\bf p}^{26}$ +33.5 ° (c 0.18, MeOH), IR (KBr) 3428, 1663, 1528, 1487, 1437, 1288; $^{1}_{\bf H}$ NMR (D₆-DMSO) δ 10.56 (1H, s), 8.71-8.57 (1H, m), 7.88-7.81 (2H, m), 7.65-
- 20 7.46 (3H, m), 4.97-4.05 (1H, m), 4.38-4.0 (3H, m), 3.88-3.52 (3H, m), 2.91-2.71 (2H, m), 2.50-2.36 (1H, m), 2.35-2.21 (1H, m), 2.10-1.94 (1H, m), 1.93-1.49 (3H, m); $^{13}{\rm C}$ NMR (D₆-DMSO) δ 173.66, 172.49, 169.97, 169.89, 164.96, 157.62, 132.35, 131.85, 128.39, 127.32,
- 25 53.81, 52.69, 40.90, 33.17, 31.60, 24.40, 24.13, 19.24; MS (ES -).

(2S) 3-Benzyloxycarbonylamino-2-phthalimidopropionic

5 acid (266). A solution of (2S) 3benzyloxycarbonylamino-2-tertbutoxycarbonylaminopropionic acid dicyclohexylamine salt (3g, 5.8mmol) in dichloromethane (200ml) was

washed four times with 1M HCl solution, dried (MgSO₄) 10 and concentrated. The resulting oil was dissolved in dry dichloromethane (35ml), cooled to 0 °C and treated with trifluoroacetic acid (35ml). This solution was stirred at 0 °C for 1.5h then evaporated to dryness.

Dichloromethane (50ml) was added to the residue then

- 15 removed under vacuum. This process repeated six times to afford a white solid. The white solid was suspended in toluene (50ml), treated with powdered phthalic anhydride (940mg, 6.35mmol) and refluxed for 18h. The resulting solution was concentrated to afford an oil
- 20 which was purified by flash chromatography (2-10-methanol/dichloromethane) to afford 266, 2.01g (94) as a white powder: 1R (KBr) 3600-2500br, 1776, 1714,

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1530, 1469, 1455, 1392, 1263, 1131, 722; ${}^{1}H$ MMR (CDCl₃) δ 7.83 (2H, m), 7.72 (2H, m), 7.29 (5H, m), 5.41 (1H, m), 5.03 (2H, s), 3.90 (2H, m); MS (ES-), 367 (M - 1).

[3S (2S)] t-Butyl 1-benzyloxycarbonyl-2-(3-

- 5 benzyloxycarbonylamino-2
 - phthalimidopropionyl)pyridazine-3-carboxylate (267). A suspension of the acid 266 (1.32g, 3.58mmol) in dry ether (37ml) was treated with phosphorus pentachloride (1.04g, 5mmol) and stirred at room temperature for 2h.
- 10 The solution was filtered to remove unreacted phosphorus pentachloride then evaporated to dryness. The residue was treated with dry toluene (25ml) then evaporated to dryness. This process was repeated several times. The resulting oil was dissolved in dry
- 15 dichloromethane (25ml), cooled to 0 °C and treated with
 a solution of (3S) t-butyl 1benzyloxycarbonylpyridazine-3-carboxylate (1.15g,
 3.58mmol) in dry dichloromethane (2ml) followed by 58

aqueous sodium bicarbonate solution (25ml). The

- 20 mixture was stirred rapidly at room temperature for 20h then diluted with ethyl acetate (100ml) and acidified to pH2 with 1M HCl. The organic phase was washed twice with dilute HCl solution then brine, dried (MgSO₄) and concentrated. The resulting oil was purified by flash
- 25 chromatography (2-20% ethyl acetate/dichloromethane
 then 10-20% methanol/dichloromethane) to afforc (267),
 1.25g (52%) as a white powder: IR (KBr) 3367, 2955,
 1722, 1517, 1455, 1387, 1369, 1251, 1153, 721; 1 HENMR
 (CDCl₃) δ 7.81 (2H, m), 7.74 (2H, m), 7.63 (1H, brs),
- 30 7.31 (10H, m), 5.46-4.76 (5H, m), 4.07-3.54 (4H, m), 2.4 (1H, m), 2.0-1.6 (3H, m), 1.40 (9H, s); MS (ES+), 671 (M + 1), 693 (M + Nai.

- 436 - (15,95) t-Butyl 1,2,3,4,7,8,9,10-octahydro-10-oxo-9-

phthalimido-6H-pyridazino[1,2-a][1,2,4]triazepine-1carboxylate (268). A solution of ester 267 (50mg, 0.074mmol) in methanol (15ml) was treated with 10% 5 palladium on carbon (50mg) and hydrogenated at room temperature and atmospheric pressure for 24h. The mixture was evacuated thoroughly to remove hydrogen then treated with 37% aqueous formaldehyde (18mg, 0.22mmol) and stirred under nitrogen for 2h. The 10 mixture was filtered, evaporated to dryness and the product purified by flash chromatography (4-100% ethyl acetate/dichloromethane) to afford 268 14.5mg (48%) as an oil: ${}^{1}H$ NMR (CDCl₃) δ 7.85 (2H, m), 7.71 (2H, m), 5.78 (1H, dd, J = 10, 5), 4.99 (1H, dd, J = 6.1, 1.5). 15 4.07 (1H, d, J = 10.6), 3.49 (1H, dd, J = 14, 5), 3.39 (1H, d, J = 10.3), 3.24 (1H, dd, J = 14, 10.2), 3.17(2H, m), 2.39 (1H, m), 1.84-1.46 (3H), 1.51 (9H, s); MS (ES+), 415 (M + 1), 437 (M + Na).

Compounds 280-283 were prepared from 212b by 20 a method similar to the method used to prepare 226e.

Compounds 284-287 were prepared by a method similar to the method used to prepare 217e.

$$Alloc \longrightarrow H$$

$$Allo$$

(35) 3-Allyloxycarbonylamino-4-oxobutyric acid tertbutyl ester O-(2,6-dichlorophenylmethyl)oxime (306a) was prepared by a similar procedure as 208a except that 2,6-dichlorophenylmethoxyamine (prepared by a similar 5 method as 306b) was used instead of semicarbazide to give 870mg (quant.) as a clear oil.

(3S) 3-Allyloxycarbonylamino-4-oxobutyric acid tertbutyl ester O-(2-(phenyl)ethyl)oxime (306b) was prepared by a similar procedure as 208a except that 2-

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(phenyl)ethoxyamine (US 5 346 911) was used instead of semicarbazide to give 395mg (quant.) as a clear oil.

[35(15,95) 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-5 [1,2-a][1,2]diazapine-1-carboxamido)-amino]-4oxobutanoicacid t-butyl ester, 0-(2,6dichlorophenylmethyl)oxime (307a) was prepared by a procedure similar to 233e except 306a was used instead

of 207a to give 23 mg(23%) of 307a as a white solid.

10 [3S(1S,9S) 3-(9-Benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazapine-1-carboxamido)-amino]-4oxobutanoic acid t-butyl ester, 0-(2-

(phenyl)ethyl)oxime (307b) was prepared by a procedure
15 similar to 233e except 306b was used instead of 207a to
give 43 mg(48%) of 307b as a white solid.

[3S(1S,9S) 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazapine-1-carboxamido)-amino]-4-

- 20 oxobutanoic acid, O-(2,6-dichlorophenylmethyl)oxime (308a) was prepared by from 307a a procedure similar to the preparation of 235e from 234e to give 15.2 mg (74)) as white solid: ¹H NMR(CD₃OD) & 0.9 (m), 1.3 (s), 1.7 (m), 1.8 (m), 2.0 (m), 2.1-2.2 (m), 2.3 (dd), 2.4-
- 25 2.5(m), 2.6(m), 2.7-2.8(m), 3.1(m), 3.3(m), 3.4-3.5(m), 4.5(m), 4.9(m), 5.1(m), 5.3(d), 5.4(s), 6.8(d), 7.2-7.5(m), 7.8(dd), 8.4(dd).

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[35(15,95) 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino-[1,2-a][1,2]diazapine-1-carboxamido)-amino]-4-oxobutanoic acid, O-(2-(phenyl)ethyl)oxime (308b) was prepared by from 307b a procedure similar to the

- 5 prepared by from 307b a procedure similar to the preparation of 235e from 234e to give 25.2 mg (68%) as white solid: 1 H NMR(CD₃OD) δ 1.2(m), 1.6-1.7(m), 2.0-2.1(m), 2.2(m), 2.3(m), 2.5(m), 2.6-2.7(dd), 2.9(t), 3.0(t), 3.1(m), 3.3-3.5(m), 4.2(t), 4.25(m), 4.5(m),
- 10 5.2(t), 5.3(t), 6.7(d), 7.1-7.2(m), 7.35(dd), 7.4(m), 7.5(m), 7.8(dd), 8.3(dd).

15

(304a) R=CH2

PCT/US96/20843

[3S(1S,9S) 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyriazino-[1,2-a][1,2]diazapine-1-carboxamido)-amino]-4-oxobutanoic acid tert-butvl ester (302).

- 5 Step A: 301 was prepared by procedure similar to 605a (Step A), except 212e was used instead of 603a to give 540 mg (34%) to give a white solid.
 Step B: 302. A solution of 301 (50.7 mg; 0.091 mmol)
 - step B: 302. A solution of 301 (50.7 mg; 0.091 mmol)
 in 2.8 ml of MeOH/HOAc/37% aq. formaldehyde (5:1:1) was
- 10 stirred at rt for 5.5 h. and the reaction was
 concentrated to 0.7 ml in vacuo. The residue was
 dissolved in 3 ml of CH₃CN and concentrated to 0.7 ml
 (3x), dissolved in toluene and concentrated to 0.7 ml
 in vacuo (2x), and concentrated to dryness.
- 15 Chromatography (flash, SiO₂, 5% isopropano1/CH₂Cl₂) gave
 302 (45.5 mg, 78%) as a white solid: 1 H NMR(DMSO-d₆) δ
 1.0-1.15(m, 2H), 1.4(s, 9H), 1.65(m, 2H), 1.9-2.1(m, 2H), 2.15-2.4(m, 3H), 2.55(m, 1H), 2.7-3.0(m, 2H), 4.3-4.6(m, 2H), 4.9(m, 1H), 5.2(m, 1H), 7.4-7.6(m, 2H),
- 20 7.8-8.0(m, 2H), 8.6(m, 1H), 8.8(m,1H), 9.4(s, 1H).

[1S,9S (2RS,3S)] 9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-methoxy-5-oxo-tetrahydro-furan-3-yl)-6H-pyridazino[1,2-a][1,2] diazapine-1-carboxamide. (304a).

- 25 Step A: A solution of 302 (90 mg; 0.18 mmol) in 10 ml of MeOH was treated with trimethylorthoformate (1ml) and p-toluene sulfonic acid hydrate (5 mg; 0.026 mmol) and the reaction was stirred for 20 h. The reaction was treated with 3 ml of aq. sat. NaHCO $_3$ and
- 30 concentrated in vacuo. The residue was taken up in

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EtOAc and washed with dilute aq. NaHCO3, dried over MgSO4 and concentrated in vacuo to give 80 mg of 303a. Step B: 303a was dissolved in 2 ml of TFA and stirred at rt for 15 min. The reaction was dissolved in CH2Cl2 and concentrated in vacuo (3x). Chromatography (flash, siO2, 1% to 3% MeOH/CH2Cl2 gave 43 mg (64%) of 304a as a white solid: ¹H NMR(CDCl3) of 1.55-1.8 (m, 2H), 1.9-2.15 (m, 4H), 2.25-2.5 (m, 2H), 2.7-3.3 (m, 4H), 3.45, 3.6(s, s, 3H), 4.4, 4.75 (2m, 1H), 4.6 (m, 1H), 4.95, 5.4 (t,d, 1H), 5.1-5.2 (m, 1H), 6.45, 7.05 (2d, 1H), 6.95 (m, 1H), 7.45 (m, 2H), 7.55 (m, 1H), 7.85 (m, 2H).

Example 11

Compounds 214e, 404-413, 415-445, 446-468, 470-491, and 493-499 were synthesized as described in 15 Example 11 and Table 7.

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Step A. Synthesis of 401. TentaGel S® NH2 resin (0.16 mmol/q, 10.0 q) was placed in a sintered glass funnel and washed with DMF (3 x 50 mL), 10% (v/v) DIEA in DMF (2 x 50 mL) and finally with DMF (4 x 50 mL). 5 Sufficient DMF was added to the resin to obtain a slurry followed by 400 (1.42 g, 2.4 mmol, prepared from (35)-3-(fluorenylmethyloxycarbonyl)-4-oxobutryic acid t-butvl ester according to A.M. Murphy et. al. J. Am. Chem. Soc., 114, 3156-3157 (1992)), 1-10 hydroxybenzotriazole hydrate (HOBT•H2O; 0.367 g, 2.4 mmol), O-benzotriazol-1-yl-N,N,N,N'-tetramethyluronium hexafluorophosphate (HBTU; 0.91 g, 2.4 mmol), and DIEA (0.55 mL, 3.2 mmol). The reaction mixture was agitated overnight at rt using a wrist arm shaker. The resin 15 was isolated on a sintered glass funnel by suction filtration and washed with DMF (3 x 50 mL). Unreacted amine groups were then capped by reacting the resin with 20% (v/v) Ac₂O/DMF (2 x 25 mL) directly in the funnel (10 min/wash). The resin was washed with DMF (3 20 \times 50 mL) and CH₂Cl₂ (3 \times 50 mL) prior to drying overnight in vacuo to yield 401 (11.0 g, quantitative

Step B. Synthesis of 402. Resin 401 (6.0 g, 0.16 mmol/g, 0.96 mmol) was swelled in a sintered glass funnel by washing with DMF (3 x 25 mL). The Fmoc protecting group was then cleaved with 25° (v/v) piperidine/DMF (25 mL) for 10 min (intermittent stirring) and then for 20 min with fresh piperidine reagent (25 mL). The resin was then washed with DMF (3 x 25 mL), followed by N-methypyrrolidone (2 x 25 mL). After transferring the resin to a 100 mL flask, N-methypyrrolidone was added to obtain a slurry followed

yield).

by 212f (0.725 g, 1.57 mmol), HOBT-H $_2$ O (C.25 g, 1.6 mmol), HBTU (0.61 g, 1.6 mmol) and DIEA (0.84 mL, 4.8 mmol). The reaction mixture was agitated overnight at rt using a wrist arm shaker. The resin work-up and 5 capping with 20% (v/v) Ac_2 O in DMF were performed as described for 401 to yield 402 (6.21 g, quantitative yield).

Step C. Synthesis of 403. This compound was prepared from resin 402 (0.24 g, 0.038 mmol) using an Advanced ChemTech 396 Multiple Peptide synthesizer. The automated cycles consisted of a resin wash with DMF (3 x 1 mL), deprotection with 25% (v/v) piperidine in DMF (1 mL) for 3 min followed by fresh reagent (1 mL) for 10 min to yield resin 403. The resin was washed with DMF (3 x 1 mL), and N-methypyrrolidone (3 x 1 mL).

Step D. Method 1. [3S(1S,9S)]-3-(6,10-Dioxo-

1,2,3,4,7,8,9,10-octahydro-9-(thiophene-3-carbonylamino)-6H-pyridazine[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (409). Resin 403 was 20 acylated with a solution of 0.4M thiophene-3-carboxylic acid and 0.4M HOBT in N-methypyrrolidone (1 mL), a solution of 0.4M HBTU in N-methylpyrrolidone (0.5 mL) and a solution of 1.6M DIEA in N-methypyrrolidone (0.35 mL) and the reaction was shaken for 2 hr at rt. The 25 acylation step was repeated. Finally, the resin was washed with DMF (3 x 1 mL), CH₂Cl₂ (3 x 1 mL) and dried in vacuo. The aldehyde was cleaved from the resin and globally deprotected by treatment with 957 TFA/5* H₂C (v/v, 1.5 mL) for 30 min at rt. After washing the

30 resin with cleavage reagent (1 mL), the combined filtrates were added to cold 1:1 EtaC:pentane (12 mL)

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and the resulting precipitate was isolated by centrifugation and decantation. The resulting pellet was dissolved in 10% CH₃CN/90% H₂O/0.1% TFA (15 mL) and lyophilized to obtain crude **409** as a white powder. The compound was purified by semi-prep RP-HPLC with a Rainin Microsorb^m C18 column (5 µ, 21.4 x 250 mm) eluting with a linear CH₃CN gradient (5% - 45%) containing 0.1% TFA (v/v) over 45 min at 12 mL/min. Fractions containing the desired product were pooled and lyophilized to provide **409** (10.8 mg, 63%).

Step D. Method 1A. Synthesis of 418. Following a similar procedure as method 1, resin 403 was acylated with 4-(1-fluorenylmethoxycarbonylamino)benzoic acid and repeated. The Fmoc group was removed as described 15 in Step C and the free amine was acetylated with 20% (v/v) Ac₂O in DMF (1 mL) and 1.6M DIEA in N-methylpyrrolidone (0.35 mL) for 2 hr at rt. The acetylation step was repeated. Cleavage of the aldehyde from the resin gave 418 (3.2 mg).

20 Step D. Method 1B. Synthesis of 447. Following a similar procedure as method 1A, rosin 403 was acylated with 0.4M 4-(1-fluorenylmethoxycarbonylamino)benzoic acid. The acylation step was repeated once. The Fmoc group was 25 removed as before and the free amine was reacted with 1M methanesulfonyl chloride in CH₂Cl₂ (0.5 mL) and 1M pyridine in CH₂Cl₂ (0.60 mL) for 4 hr at rt. Cleavage of the aldehyde from the resin gave 447 (10.0 mg).

Step D. Method 2. Synthesis of 214e. Following 30 a similar procedure as method 1, resin 403 was advlated

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with 0.5M benzoyl chloride in N-methypyrrolidone (1 mL) and 1.6M DIEA in N-methypyrrolidone (0.35 mL) for 2 hr at rt. The acylation step was repeated. Cleavage of the aldehyde from the resin gave 214e (5.1 mg, 30%).

- 5 Step D. Method 3. Synthesis of 427. Following a similar procedure as method 1, resin 403 was reacted with 1.0M benzenesulfonyl chloride in CH₂Cl₂ (0.5 mL) and 1M pyridine in CH₂Cl₂ (0.60 mL) for 4 hr at rt. The reaction was repeated. Cleavage of the aldehyde 10 from the resin gave 427 (7.2 mg, 40%).
- Step D. Method 4. Synthesis of 420. Following a similar procedure as method 1, resin 403 was reacted with 0.5M methylisocyanate in N-methypyrrolidone (1 mL) and 1.6M DIEA in N-methypyrrolidone (0.35 mL) for 2 hr at rt. The reaction was repeated. Cleavage of the aldehyde from the resin gave 420 (8.3 mg, 55%).
- Step D. Method 5. Synthesis of 445. Following a similar procedure at method 1, resin 403 was acylated with 0.27M imidazole-2-carboxylic acid (1 mL) in 2:1 20 DMF:H₂O (with 1 eq. DIEA) and IM 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDC) in 2:1 N-methypyrrolidone/H₂O (0.35 mL) for 3 hr at rt. Cleavage of the aldehyde from the resin gave 445 (9.5 mg).

25 Analytical HPLC methods:

(1) Waters DeltaPak C18, 300A (5 μ , 3.9 x 150 mm). Linear CH₃CN gradient (5 $\hat{\gamma}$ - 45 $\hat{\gamma}$) containing 0.1° TFA (ν/ν) over 14 min at 1 mL/min.

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- (2) Waters DeltaPak C10, 300A (5 μ , 3.9 x 150 mm). Linear CH₃CN gradient (0% 25%) containing 0.1% TFA (v/v) over 14 min at 1 mL/min.
- (3) Waters DeltaPak C18, 300A (5μ, 3.9 x 150 mm).
- 5 Isocratic elution with 0.1% TFA/water (v/v) at 1 mL/min.
 - (4) Waters DeltaPak C18, 300A (5μ , 3.9 x 150 mm). Linear CH₃CN gradient (0% - 30%) containing 0.1% TFA (v/v) over 14 min at 1 mL/min.
- 10 (5) Waters DeltaPak C18, 300A (5µ, 3.9 x 150 mm). Linear CH₃CN gradient (0% - 35%) containing 0.1% TFA (v/v) over 14 min at 1 mL/min.

Cmpd.	Structure	MF	мм	HPLC RT	MS (M+H) +	Syn. Method
214e		C21H24N4O7	444.45	6.67 (2)	445	7
404		C22H26N4O7	458.48	6.66 (2) 978	459	2
405		C22H26N4O8	474.47	8.2 (1) 98%	475	5

Cmpd.	Structure	MF	M	HPLC RT	MS	Syn.
				min	+ (H+W)	Method
406	CONTRACTOR OF THE CONTRACTOR O	C21H23C1N407	478.89	6.33 (1) 98%	479	2
407	E H	C25H26N4O7	494.51	9.90 (1)	495	7
408		C25H26N4O7	494.51	9.0 (1)	495	7
409		C27H28N4O7	520.55	11.14 (1) 98%	521	0

Cmpd.	Structure	MF	ММ	HPLC RT	MS + (H+H)	MS Syn.
410	HOW WE WANTED	C19H22N407S	450.47	4.87 (1) 98%	451	1
411		C24H25N5O7	495.50	10.7 (1)	496	1
412	io iii o	C24H25N5O7	495.50	8.57 (1) 98%	496	
413	O N N N N N N N N N N N N N N N N N N N	C18H24N4O7	408.41	7.21 (2) 98%	409	

Cmpd.	Structure	[1 X	3M	HPLC RT	MS	Syn.
				min	+ (H+H)	(M+H) + Method
415		C22H24N4O9	488.46	7.58 (1)	489	н
416		C21H23C1N407	478.89	9.66 (1)	479	r-d
417		C24H30N4010	534.53	8.12 (1) 535	535	1

Structure MF MW		Æ	>	HPLC RT	WS W	Syn.
		-		uım	+ (H+W)	Method
HN ON C23HZ7N508	C23H27N5O8		501.50	5.93 (1)	502	1A
S S S S S S S S S S S S S S S S S S S	C16H22N408		398,38	6.84 (2)	399	7
H H CIGHZ3NSO7	C16H23N5O7		397.39	5.25 (2)	398	4

T C	0 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	M	204	HPLC RT	MS	Syn.
24	Octaconto	7115	u U	min	+ (H+W)	Method
421		C16H24N408S	432.46	7.13 (2)	433	m
422	N H O H OH	C21H28N6O7	476.49	6.89 (1) 98%	477	-
423	O N O O O O O O O O O O O O O O O O O O	C20H25N507S	479.52	5.62 (1) 98%	480	11
424		C19H23N5O8	449.42	6.28 (1)	450	1

Cmpd.	Structure	M	ММ	HPLC RT min	MS (M+H)+	MS Syn. (M+H)+ Method
425	LOCAL OF THE STATE	C25H26N408	510.51	8.25 (1) 98%	511	п
426	H COH	C21H30N4O7	450.50	8.0 (1) 98%	451	7
427	O NI NI O	C20 H24N 408S	480.50	7.87 (1)	481	м

Стрд.	Structure	MF	ММ	HPLC RT min	MS (M+H) +	Syn. Method
428	O O O O O O O O O O O O O O O O O O O	C16H25N508S	447.47	5.13 (1)	448	т
429	H, N, H	C14H20N4O6	340.34	3.19 (3) 98%	341	
430	NH O	C23H27N5O8	501.50	5.53 (1) 988	502	1.8
431	HO H	C21H25N5O7	459.46	6.66 (2) 98%	460	-

C	44,000	i k	200	HPLC RT	MS	Syn.
))))	111	17100	min	+ (H+H)	Method
432	D I I I I I I I I I I I I I I I I I I I	C21H23N7O7	485.46	5.59 (1) 98%	486	1
433	To Not Not Not Not Not Not Not Not Not No	C24H27N5O7	497.51	11.07 (1)	498	et
434	A TOO MANAGEMENT OF THE PROPERTY OF THE PROPER	C22H24N6O7	484.47	4.43 (1) 98%	485	н
435	O N N N N N N N N N N N N N N N N N N N	C24H25N5O7	495.50	5.10 (1)	496	1

Cmpd.	Structure	i. X	Met	HPLC RT	MS	Syn.
2	2000	ME	MILI	min	+ (H+H)	Method
436	HO THE PART OF THE	C24H25N5O7	495.50	8.20 (4) 98%	496	
437	N N N O N N N N N N N N N N N N N N N N	C25H27N5O8	525.52	12.78 (5) 98%	526	
438		C24H25N5O7	495.50	4.85 (1) 98%	496	п
439	OF ZZ OO ZZ	C24H25N5O7	495.50	8.70 (5) 98%	496	1

Cmpd.	Structure	Σ	MW	HPLC RT	MS	Syn.
				min	(M+H) +	(M+H) + Method
440		C25H27N5O7	509.52	9.96 (5)	510	1
441	Po No	C27H31N5O7	537.58	6.15 (1) 988	538	1
442	H H H H S S	C21H22N407S2	506.56	10.10 (1)	507	1

Cmpd.	Structure	MF	MW	HPLC RT	MS	Syn.
				min	(M+H) +	(M+H) + Method
443	## ## ## ## ## ## ## ## ## ## ## ## ##	C27H28N4O8	536.55	13.12 (1) 98%	537	T.
444	O ZI O O ZI O O ZI O	C21H22C12N407	513.34	9.96 (5) 988	510	-
445	HO H HO N H HAVE	C18H22N6O7	434.41	5.72 (1) 98%	435	ဟ

				HDIC DI	MC	41.0
Cmpd.	Structure	MF	MM	101 707 111	CIT	. n.k.c
				min	+ (H+H)	Method
446	O NI O NI O NI O NI	C17H20N607S	452.45	5.00 (1) 983	453	Н
447	ST CO	C22H27N509S	537.55	6.32 (1) 98§	538	1.18
448	A COMPANY OF THE COMP	C24H29N5O8	515.53	6.36 (1) 98%	516	I.A.

Structure	MF	MW	HPLC RT min	MS (M+H)+	MS Syn. (M+H)+ Method
O Z Z PO NI O Z Z PO NI O Z Z PO	C25H26N4O8	510.51	13.86 (1)	511	1
DE STEED OF THE PERSON OF THE	C23H27N5C8	501.50	6.10 (1)	502	LA
O NH	C22H26N4O8	474.47	8.02 (1) 988	475	2

				EG CIGH	o _M	1
Structure		MF	MM	THE TO LET	CE .	· uke
	- 1			min	+ (H+W)	Method
		C22H2 6N408	474.47	7.77 (1)	475	2
S S S S S S S S S S S S S S S S S S S		C23H24N407S	500.53	11.11 (1) 98%	501	2
O Z Z O Z Z O Z Z Z O Z Z Z O Z Z Z O Z Z Z O Z Z Z O Z Z Z O Z Z Z Z Z O Z		C20H23N5O7	445.44	6.24 (2)	446	7
NH NH NH		C21H23C1N407	478.89	9.45 (1)	479	2

Cmpd.	Structure	MF	MM	HPLC RT	MS	Syn.
				min	+ (H+W)	Method
456	ST S	C21H24N4O8	460.45	5.58 (1)	(M+Na) 483	н
457	NAT OF THE PROPERTY OF THE PRO	C28H28N4O10	580.56	10.42 (1)	(M+Na) 603	Н
458	O Z Z Z O O Z Z Z O O	C21H22F2N407	480.43	8.65 (1) 98%	481.1	_

Cmpd.	Structure	2	34	HPLC RT	MS	Syn.
		111	LIM.	min	+ (H+W)	Method
459		C21H22C1FN407	496.88	10.11 (1) 98%	498.3	
460	H ₃ C ₂ C ₃	C22H26N409S	522,54	6.16 (1)	523.6	М
461	O NH ON NH ON NH ON NH	C21H23FN407	462.44	7.41 (1)	463.3	1

					-	
Cmpd.	Structure	ΜŁ	MΜ	HPLC RT	MS	Syn.
				min	+ (H+W)	(M+H) + Method
462	ST S	C21H23FN4O7	462.44	7.71 (1)	463.3	П
463	O N N N N N N N N N N N N N N N N N N N	C21H23FN407	462.44	7.64 (1)	464	1
464	C C C C C C C C C C C C C C C C C C C	C21H22C12N407	513.34	11.59 (1) 988	414.5	1

Cmpd.	Structure	Μ	ММ	HPLC RT	MS (M+H)+	MS Syn. (M+H)+ Method
	HO HO JHO	C22H25C1N407	492.92	9.65 (1)	493.9	1
Į.	HJC O W O N	C22H25C1N4O7	492.92	9.63 (1)	493.9	1
		C23H24N4O8	484.47	9.73 (1)	485.8	1

Cmpd.	Structure	MF	MW	HPLC RT	MS	Syn.
				min	+ (H+H)	Method
468	S	C26H26F3N507S	609.59	14.84 (1)	609.7	1
470	H. C. O. C.	C23H29N5O7	487.52	4.57 (1) 98%	489.5	1
471	H ₃ C _N H ₃	C23H29N5O7	487.52	5.74 (1)	488.2	1

Cmpd.	Structure	X.	32	HPLC RT	MS	Syn.
				min	+ (H+W)	Method
472	O Z Z O NI	C22H25N5O7	471.47	4.00 (1)	474	п
473		C23H26N4O9	502.49	7.65 (1)	503.6	erd.
474	P H C H C H C H C H C H C H C H C H C H	C23H26N408	486.49	7.16 (1) 98%	488.1	1

E C	orutourto	1 2	S.	HPLC RT	MS	Syn.
· polimo	מרומות	INE	M.	min	+ (H+W)	(M+H) + Method
475	HO N H HY	C23H25N5O7	483.49	9.77 (1) 978	485.1	
476	HO H	C22H26N4O8	474.47	5.25 (1) 98%	475.8	
477	N. N	С2 6Н33N5О9	559.58	4.76 (1) 95%	561.8	- 7

			-			
Cmpd.	Structure	Σ	MM	HPLC RT	MS	Syn.
				min	+ (H+H)	Method
478	H H H H H H H H H H H H H H H H H H H	C21H25N509S	523.53	5.25 (1)	524.3	
479	HO LA COMPANY	C22H26N408	474.47	5.35 (1) 98%	475.8	
480	HO H	C25H30N6O9	558.55	5.11 (1)	559.3	1A

	·			HPLC RT	MS	Svn.
cimpa.	erinciale	Į.	MM	min	+ (H+H)	Method
481	HO NH O NH O NH	C21H24CIN5O7	493.9	7.10 (1)	495.1	1
482	O NH O NH O NH	C21H23C12N507	528.4	9.05 (1)	529.8	-4
483		C28H29N5O8	563.57	10.01 (1)	565.6	1,2

Cmpd.	Structure	Ľ.	MM	HPLC RT	MS	Syn.
				min	+ (H+H)	(M+II) + Method
484	H _{GC}	C25H31N5O8	529.55	7.88 (1) 98%	531	1,2
485	Hychin Hoom	C24H29N5O8	515.53	7.00 (1)	517.6	1,2
486		C29H31N5O8	577.60	10.43 (1)	579.4	1,2

Structure	MF	MM	HPLC RT min	MS + (H+H)	Syn. Method
H, C M M M M M M M M M M M M M M M M M M	C2 6H33N5O8	543.58	9.30 (1)	545.7	1,2
H _F	C25H31N5O8	529.55	8.13 (1) 98%	531.1	1,2
H-CH H-CH H-CH H-CH H-CH H-CH H-CH H-CH	C23H28N6O8	516.52	5.89 (1) 98%	517.8	1,4
T Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	C23H27N509	517.50	7.27 (1)	(M+Na) 540.8	1,2

Cmpd.	Structure	Α	30	HPLC RT	WS	Syn.
				min	+ (H+H)	Method
491		C28H28N4O9	564.56	12.9 (1) 98%	565.3	-1
493	H _{5CO} F C C C C C C C C C C C C C C C C C C	C22H25FN408	492.46	8.31 (1) 98%	493.9	1
494	HO N N N N N N N N N N N N N N N N N N N	C23H26N4O7	470.49	9.34 (1) 98%	471.2	2

Cmpd	Structure	ía X	30	HPLC RT	MS	Syn.
,	h 123332		110	min	+ (H+H)	Method
495	O ZIZOO	C22H26N4O7	458.48	7.24 (1)	459.9	7
496	HO NO	C22II26N408	474.47	9.47 (1)	475.7	2
497	HAND OF H	C22H25C1N408	508.92	9.58 (1) 98%	509.5	
498	O Z Z PO O Z PO D O Z PO D O Z PO	C21H23C1N408	494.89	7.18 (1)	495.1	-

npd.	Structure	Ē	32	HPLC RT	MS	Syn.
				min	(M+H) + Method	Method
661	Hot Strong	C28H30N4O8	550.57	13.27 (1) 98%	552	

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Example 12

Compounds 605a-j, 605m-q, 605s, 605t, and 605v were synthesized as described below.

Compound no.	R ₂	R ₅
600a/103	Н	CH ₃
600ъ	Н	CH ₂ Ph
600c	CH ₂	CH ₂ Ph

5

(3S)-2-0xo-3-tert-butoxycarbonylamino-2,3,4,5tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl 10 ester (600a/103).

Step A. (2S)-2-tert-Butoxycarbonylamino-3-(2nitrophenyl-amino)-propionic acid. (2S)-2-tertButoxycarbonylamino-3-aminopropionic acid (10 g,
49 mmol), 2-fluoronitrobenzene (5.7 ml, 54 mmol), and
15 NaHCO3 (6.25 g, 98 mmol) was taken into 130 ml of DMF

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and heated at 80 °C for 18 h. The reaction was evaporated in vacuo to give a viscous orange residue that was dissolved in 300 ml of $\rm H_2O$ and extracted with Et₂O (3 x 150 ml). The aq. solution was acidified to 5 pH 5 with 10% NaHSO₄ and extracted with EtOAC (3 x 250 ml). The combined extracts were dried over anhydrous Na₂SO₄, filtered, and evaporated to give 12.64 g (83%) of the title compound as an orange amorphous solid: $^{1}{\rm H}$ NMR (CD₃OD) δ 8.15-8.10 (1H,d), 7.54-7.48 (1H,t), 7.13-7.08 (1H,d), 6.73-6.65 (1H,t), 4.45-4.35 (1H,m), 3.9-3.8 (1H,dd), 3.65-3.55 (1H,dd), 1.45 (9H,s).

<u>Step B.</u> (2S)-2-tert-Butoxycarbonylamino-3-(2-aminophenyl-amino)-propionic acid. A mixture of (2S)-2-tert-Butoxycarbonylamino-3-(2-

- 15 nitrophenylamino)propionic acid (12.65 g, 40.5 mmol) and 0.5 g of 10% Pd/C in 100 ml of MeOH under hydrogen at 1 atmosphere was stirred for 4 h. The solution was filtered through Celite 545 and the filtrate evaporated in vacuo to afford the 11.95 g of the title compound in quantitative yield as a dark brown solid that was used
- without purification: 1 H NMR (CD₃OD) δ 6.75-6.70 (3H, m), 6.65-6.58 (1H, m), 4.35-4.3 1H, m), 3.6-3.38 (2H, m), 1.45 (9H, s).

Step C. (3S)-2-0xo-3-tert-Butoxycarbonylamino-1,3,4,5-

25 tetrahydro-1H-1,5-benzodiazepine. 1-(3Dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride
(8.54 g, 44.5 mmol) was added to a cooled (0 °C)
solution of (25)-2-tert-butoxycarbonylamino-3-(2-

aminophenylamino)propionic acid (11.95 g, 40.5 mmol) in 30 100 ml of DMF and stirred for 18 h. The reaction was poured into 700 ml of EtOAc and washed four rimes with

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100 ml of H₂O. The organic layer was dried over anhydrous Na₂SO₄, filtered, and evaporated to give a brown solid that was purified by flash chromatography eluting with 3:7 EtOAc/hexane to give 8 g (71%) of the 5 title compound: ^{1}H NMR (CDCl₃) δ 7.78 (1H, s), 7.02-6.95 (1H, m), 6.88-6.82 (1H, m), 6.82-6.78 (1H, m), 6.75-6.70 (1H, m), 5.8-5.7 (1H, d), 4.55-4.45 (1H, m), 3.95 (1H, s), 3.9-3.82 (1H, m), 3.48-3.40 (1H, m), 1.45 (9H,s).

- Step D. (3S)-2-Oxo-3-tert-butoxycarbonylamino-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl ester (600a/103). A 1.0 M solution of lithium bis(trimethylsilyl)amide (3.4 ml, 3.4 mmol) in THF was added dropwise to a -78 °C solution of (3S)-2-oxo-3-
- 15 tert-butoxycarbonylamino-2,3,4,5-tetrahydro-1H-1,5benzodiazepine (0.94 g, 3.38 mmol) in 20 ml of
 anhydrous THF and stirred for 30 min. Methyl
 bromoacetate (0.44 ml, 4 mmol) was added dropwise to
 the reaction mixture then warmed to RT. The reaction
 20 was diluted with 100 ml of EtoAc and washed with 0.3N
 KHSO₄ (50 ml), H₂O (2 x 50 ml), and brine. The
 combined organics were dried over anhydrous Na₂SO₄,
 filtered, and evaporated to afforded a gum that was
- 25 EtOAc/Hex. to give 0.98 g (83%) of the title compound as a white solid. ¹H NMR (CDCl₃) δ 7.15-7.07 (2H, m), 6.98-6.94 (1H, m), 6.88-6.84 (1H, d), 5.62-5.55 (1H, d), 4.71-4.65 (1H, d), 4.65-4.6 (1H, m), 4.33-4.27 (1H, d), 3.96-3.90 (1H, m), 3.78 (3H, s), 3.44-3.37 (1H, m),

purified by flash chromatography eluting with 3:7

30 1.4 (9H, s).

(3S)-2-0xo-3-tert-butoxycarbonylamino-2,3,4,5tetrahydro-1H-1,5-benzodiazepine-1-acetic acid benzyl ester (600b). Prepared by a similar method described for the preparation of 600a/103 (Step D), except benzyl 5 bromoacetate was used instead of methyl bromoacetate to give 600b in quantitative yield.

(3S)-2-0xo3-tert-butoxycarbonylamino-2,3,4,5tetrahydro-7,9-dimethyl-1H-1,5-benzodiazepine-1-acetic acid benzyl ester (600c).

- Step A. (25)-2-tert-Butoxycarbonylamino-3-(2-nitro-3,5-dimethylphenylamino)-propionic acid. Prepared by a method similar as described for 600a/103 (Step A), except 2-fluoro-4,6-dimethyl-nitrobenzene was used instead of 2-fluoronitrobenzene to give the desired compound in 93% yield.
- Step B. (2S)-2-tert-Butoxycarbonylamino-3-(2-amino-3,5-dimethylphenyl-amino)-propionic acid. (2S)-2-tert-Butoxycarbonylamino-3-(2-nitro-3,5-dimethylphenylamino)propionic acid was converted to the title 20 compound in quantitive yield as described in the prepartation of 600a/103 (Step B).

Step C. 2-0xo-(3S)-3-tert-butoxycarbonylamino-2,3,4,5-

tetrahydro-7,9-dimethyl-1H-1,5-benzodiazepine. A 0 °C solution of (2S)-2-tert-butoxycarbonylamino-3-(2-amino-25 3,5-dimethylphenyl-amino)-propionic acid (763 mg, 2.36 mmol) and N-methylmorpholine (483 mg, 4.78 mmol) in 60 ml of anhydrous THF was treated dropwise with isobutylchloroformate (352 mg, 2.5 mmol). The reaction was stirred for 2 h at 0 °C, at RT for 1h and poured over EtOAc. The mixture was washed with aq. 5- NAHSO4,

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sat. aq. NaHCO3, and sat. aq. NaCl, dried over NaSO4, and concentrated in vacuo. Chromatography (flash, SiO2, 10% to 25% to 50 % EtOAc/CH2Cl2) gave 490 mg (68%) of the desired product.

5 Step D. (3S)-2-Oxo-3-tert-butoxycarbonylamino-2,3,4,5-tetrahydro-7,9-dimethyl-1H-1,5-benzodiazepine-1-acetic acid benzyl ester (600c). (2S)-2-tert-Butoxycarbonylamino-3-(2-amino-3,5-dimethylphenyl-amino)-propionic acid was converted to 600c, 75% by a 10 similar method for the preparation of 600b.

(3s)-2-0xo-3-benzoylamino-5-(3-phenylpropionyl)2,3,4,5-tetrahydro-1H-1,5-benzo diazepine-1-acetic acid
methyl ester (602a).

Step A. Anhydrous HCl was bubbled into a solution of 15 (3S)-2-oxo-3-tert-butoxycarbonylamino-2,3,4,5tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl ester (600a/103, 4.0 g, 11.4 mmol) in 20 ml of CH₂Cl₂ for 20 min then stirred for 1 h at RT. The reaction was evaporated to give (3S)-2-oxo-3-amino-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl ester hydrochloride as a white solid.

- Step B. The white solid was dissolved in 70 ml of DMF and benzoic acid (1.5 g, 12.3 mmol) was added. The reaction was cooled in a ice/H₂O bath and treated with 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (2.4 g, 12.5 mmol), 1-hydroxybenzotriazole (1.7 g, 12.6 mmol) and
- diisopropylethylamine (3.0g, 23.2 mmol). The reaction was stirred for 18 h at RT under nitrogen atmosphere and poured onto $\rm H_2O$. The aq. mixture was extracted with EtOAc (2x). The combined organic layers were washed with aq. 0.5 N NaHSO4, $\rm H_2O$, sat. aq. NaHCO3, $\rm H_2O$
- 15 and sat. aq. NaCl, dried over MgSO₄ and concentrated in vacuo. Chromatography (flash, SiO₂, 10% to 30% EtOAc/CH₂Cl₂) gave 3.4 g (85%) of (3S)-2-oxo-3-(benzoylamino)-2,3,4,5-tetrahydro-1H-1,5benzodiazepine-1-acetic acid methyl ester as a white
- 20 solid.

Step C. Method A. (38)-2-0xo-3-benzoylamino-5-(3phenylpropionyl)-2,3,4,5-tetrahydro-1H-1,5benzodiazepine-1-acetic acid methyl ester (602a). A
solution of (38)-2-0xo-3-(benzoylamino)-2,3,4,525 tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl

- ester (200 mg, 0.57 mmol) in CH₂Cl₂(10 ml) was treated with triethylamine (119 mg, 1.13 mmol) and 3-phenylpropionyl chloride (114 mg, 0.68 mmol). The reaction was stirred at RT for 30 min and diluted with
- 30 CH2Cl2. The solution was washed with aq. 10% HCl, sat. aq. NaHCC3 and sat. aq. NaCl, dried over Na_2SO_4 and

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concentrated in vacuo to give 240 mg (87%) of ${\bf 602a}$ as a white foam.

Step C. Method B. (3S)-2-Oxo-3-benzoylamino-5-acetoacety1-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-5 acetic acid benzyl ester (602g). A 0 °C solution of (3S)-2-oxo-3-(benzoylamino)-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid benzyl ester (600b) (465 mg, 1.10 mmol) in CH₂Cl₂ (5 ml) was treated with acetoacetic acid in 1 ml of CH₂Cl₂ followed by slow addition of 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (431 mg, 2.2 mmol) in 2 ml of CH₂Cl₂ under N₂ atmosphere. After 15 min the reaction was poured onto EtOAc, washed with aq. 5 % NaHSO₄, dried over Na₂SO₄ and concentrated in vacuo.
15 Chromatography (flash, SiO₂, 0% to 10% to 25% MeOH/CH₂Cl₂) gave 580 mg of (3S)-2-oxo-3-(benzoylamino)-5-acetoacety1-2,3,4,5-tetrahydro-1H-1,5-benzodiazening-lacetic acid benzul aceta aceta.

- (benzoylamino)-5-acetoacety1-2,3,4,5-tetrahydro-1*H*-1,5-benzodiazepine-1-acetic acid benzyl ester as a white solid.
- 20 <u>Step C. Method C.</u> (3S)-2-0xo-3-benzoylamino-5-methoxycarbonyl-2,3,4,5-tetrahydro-1H-1,5-benzo diazepine-1-acetic acid benzyl ester (602j). A vigorously-stirred, 0 °C solution of (3S)-2-0xo-3-(benzoylamino)-2,3,4,5-tetrahydro-1H-1,5-
 - 5 benzodiazepine-1-acetic acid benzyl ester (600b) (461 mg, 1.07 mmol) in THF (5 ml) and sat. aq. NaHCO₃ (2.5 ml) was treated with a THF solution (0.35 ml) of methyl chloroformate (151 mg, 1.6 mmol) and the reaction was stirred for 45 min at RT. The reaction
- 30 was poured onto CH_2Cl_2 and washed with H_2O , dried over Na_2SO_4 and concentrated in vacuo. Chromatography

(flash, ${\rm SiO_2}$, 0% to 10% MeOH/CH $_2{\rm Cl_2}$) gave 525 mg of 602 $_{\rm J}$ as a white solid.

- Step C. Method D. (3S)-2-Oxo-3-benzoylamino-5-benzylaminocarbonyl-2,3,4,5-tetrahydro-1H-1,5-
- 5 benzodiazepine-1-acetic acid methyl ester (602p). A solution of 600a/103 (400 mg, 1.1mmol) and benzylisocyanate (166 mg, 1.2mmol) in 10 ml of CH_2Cl_2 and 10 ml of DMF and heated at 80 °C for 3 days. The reaction was cooled to RT poured onto H_2O and extracted
- 10 with EtOAc (2x). The combined organic layers were washed with H₂O (4x) and sat. aq. NaCl, dried over MgSO₄ and concentrated in vacuo. Chromatography (flash, SiO₂, 50% to 80% EtOAc/hexane) gave 440 mg (80%) of 602p as a white solid.
- 15 Step C. Method E. (3S) 2-0xo-3-benzylamino-5-(3-phenylpropionyl)-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl ester (602v). A solution of (3S) 2-0xo-3-amino-5-(3-phenylpropionyl)-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl ester hydrochloride (560 mg, 1.34 mmol), benzaldebyda (146 mg, 1.34 mmol), and sodium acetyte
- benzaldehyde (146 mg, 1.34 mmol) and sodium acetate (220 mg, 2.68 mmol) in methanol (20 ml) was treated with 4Å sieves (2 g) and NaCNBH3 (168 mg, 2.68 mmol). The reaction was stirred for 2.5 h, acidified with 10
- 25 aq. HCl to pH 2 and washed with Et₂O (2x75 ml). The organic layers were concentrated in vacuo to give an oil. Chromatography (flash, SiO₂, 0 to 35° EtOAc/CH₂Cl₂° gave 250 mg (40%) of 602v as a clear oil.

Step D. Method A. (3S)-2-Oxo-3-benzoylamino-5-(3-

30 phenylpropionyl)-2,3,4,5-tetrahydro-1H-1,5-

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benzodiazepine-1-acetic acid (603a). (3S)-2-0xo-3-benzoylamino-5-(3-phenylpropionyl)-2,3,4,5-tetrahydro-1H-1,5-benzo diazepine-1-acetic acid methyl ester (602a; 1.25 g, 2.57 mmol) was dissolved in 11 ml of 5 THF, MeOH and H_2 O (5:5:1) and treated with LiOH- H_2 O (42 mg, 0.62 mmol) stirred at RT for 64 h. The reaction was concentrated *in vacuo*, diluted with H_2 O and acidified with aq. 1N HCl to give 230 mg of 603a as a white solid.

10 Step D. Method B. (3S) 2-0xo-3-benzoylamino-5-acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid (603d). A mixture of (3S)-2-0xo-3-(benzoylamino)-5-acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid benzyl ester (602d; 510 mg, 1.08 mmol) and 15 5% Pd/C (250 mg) in MeOH (10 ml) stirred under H₂ (1 atm) for 0.5h. The reaction was filtered and concentrated in vacuo 410 mg of 603d as a white solid.

The compounds of Table 8 were prepared as described in Table 9, using the methods of Example 12.

20 Table 8

Compound R ₂		R ₃	R ₄	R ₅
602b	Н	PhCH ₂ C(O)	PhC (O)	CH ₂ Ph
602c	Н	PhC (0)	PhC(O)	CH ₂ Fh
602d H 602e H		CH3C(0)	PhC (O)	CH ₂ Ph
		CH3OCH2C(O)	PhC (O)	CH ₂ Ph
602f	Н	(CH ₃) ₂ CHCH ₂ C(O)	PhC (O)	CH ₂ Ph
602g	Н	CH3C (0) CH2C (0)	PhC(O)	CH, Ph

	Compound no.	R ₂	R ₃	R ₄	R ₅
	602h	Н	CH3OC (O) C (O)	PhC (0)	CH ₂ Ph
	602i	н	CH ₃ C (0) C (0)	PhC(O)	CH ₂ Ph
	602j	Н	CH3OC(0)	PhC (0)	CH ₂ Ph
	602k	Н	CH ₃ C (0)	Вос	CH ₂ Ph
5	6021	СНЗ	CH ₃ C(0)	Вос	CH ₂ Ph
	602m	Н	CH3S (O2)	PhC(O)	CH ₃
	602p	Н	PhCH ₂ NHC (O)	PhC(O)	CH ₃
	602q	Н	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	PhC(O)	CH ₂ Ph
	602r	Н	PhCH ₂ CH ₂ C (O)	PhCH ₂ CH ₂ C(O)	CH ₂ Ph
10	602s	Н	4-pyridylCH ₂ C(O)	PhC (0)	CH ₂ Ph

Table 9

No.	Starting material	R₃X	Step C method/ (% yield)	Step D method/ (% yield)	
603b	600Ь	PhCH ₂ C(O)Cl	A (98)	B (89)	
603c	600ь	PhC (0) Cl	A (quant.)	B (quant.)	
603d	600b	CH3C(0)Cl	A (quant.)	B (quant.)	
603e	600ь	CH3OCH2C(0)Cl	A (59)	B (quant.)	
603f	600ъ	(CH ₃) ₂ CHCH ₂ C(0)Cl	A (88)	B (95)	
603g	600ь	CH ₃ C(0)CH ₂ CO ₂ H	B (quant.)	B (quant.)	
603h	600b	CH30C(0)C(0)C1	A (96)	B (quant.)	
603i	600ь	CH ₃ C(0)CO ₂ H	B (87)	B (94)	
603ј	600ь	CH3OC(0)Cl	C (quant.)	B (quant.)	

No.	Starting material	R₃X	Step C method/ (% yield)	Step D method/ (% yield)
603k	600Ъ	СН ₃ С(О)С1	A, Step C only (quant.)	not run
6031	600c	CH3C(0)Cl	A, Step C only (quant.)	not run
603m	600a/103	CH ₃ SO ₃ Cl, NEt ₃ instead of pyridine and THF instead of CH ₂ Cl ₂	A (76)	A (92)
603p	600a/103	PhCH ₂ C=N=O	D (80)	A (86)
603q	600b	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	C (83)	B (71)
603r	600a/103	PhCH ₂ CH ₂ C(0)Cl	A	
603s	600Ъ	4-pyridylCH ₂ CO ₂ H	B (90)	B (98)

5

Boc-N R₂

1) Step C. R₃X

R₂

2) Step A. HCl

3) Step C. R₄X

R₄

R₄

R₇

R₂

600

R₄

Step D

R₄

R₂

R₃

R₄

R₄

R₂

602

The compounds of Table 10 were prepared as described in Table 11 using the methods of Example 12.

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Table 10

Compound R ₂		R ₃	R ₄	R ₅
602n	Н	CH ₃ C (O)	Naphthylene-2-C(O)	CH ₂ Ph
6020	CH ₃	CH3C(0)	PhC(O)	CH ₂ Ph
602t	Н	3-CH ₃ PhCH ₂ C(0)	PhC (0)	CH ₂ Ph
602u	Н	CH3C(0)	Fmoc	CH ₂ Ph
602v	Н	PhCH2CH2CO	PhCH ₂	CH ₂

Table 11

10	No.	Starting material	1) Step C. R ₃ X method (% yield)	3) Step C R ₄ X method (% yield)	Step D method (% yield)
	603n	602k	CH ₃ C(O)Cl A (quant.)	naphthylen e- 2-C(O)Cl A (70)	B(quant.)
	603o	6021	CH ₃ C(O)Cl A (quant.)	PhC(C)Cl A (73)	B(quant.)
	603t	602k	3- CH ₃ PhCH ₂ C(0)C1 A (quant.)	PhC(O)Cl A (93)	B (95)
	603u	602k	CH ₃ C(O)Cl A (quant.)	Fmoc-C1 C (82)	C (98)
15	603v	600a/103	2 - 2 - 1 - 7	PhCHO E(40)	A (95)

 $\begin{tabular}{ll} The compounds of Table 12 were prepared by \\ the methods described below. \end{tabular}$

Table 12

compound no.	R ₂	R ₃	R ₄
605a	н	PhCH ₂ CH ₂ C(O)	PhC (0)
605b	Н	PhCH ₂ C(O)	PhC (0)
605c	Н	PhC (0)	PhC (0)
605d	Н	CH ₃ C (O)	PhC (0)
605e	Н	CH3OCH2C(O)	PhC (0)
605f	Н	(CH ₃) ₂ CHCH ₂ C(O)	PhC(O)
605g	Н	CH ₃ C (0) CH ₂ C (0)	PhC (0)
605h	Н	CH3OC (0) C (0)	PhC (O)
605i	Н	CH ₃ C (0) C (0)	PhC (O)
605j	Н	CH ₃ OC (O)	PhC (0)

10

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compound no.	R ₂	R ₃	R ₄
605m	Н	CH3SO3	PhC(O)
605n	н	CH ₃ C(O)	Naphthy1-2-C(0)
605o	CH ₃	CH3C(O)	PhC (O)
605p	Н	PhCH ₂ NHC (0)	PhC (0)
605q	Н	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	PhC (O)
605s	Н	4- pyridylCH ₂ C(O)	PhC(O)
605t	Н	3-CH ₃ PhCH ₂ C(O)	PhC (0)
605v	Н	PhCH ₂ CH ₂ C(O)	PhCH ₂

(3S) -3-[(3S) -2-0xo-3-benzoylamino-5-(3-

5

phenylpropionyl)-2,3,4,5-tetrahydro-1H-1,5benzodiazepin-1-acetylamino]4-oxo-butyric acid (605a).

<u>Step A.</u> (35)-3-(1-Fluorenylmethyloxycarbonylamino)-4-oxobutyric acid tert-butyl oster semicarbazone (210 mg, 0.45 mol, Prepared in a similar manner to the

- 15 benzyloxycarbonyl analog in Graybill et al., Int. J.
 Protein Res. , 44, pp. 173-82 (1994).) was dissolved in
 10 nl of DMF and 2 ml of diethylamine and stirred for 2
 h. The reaction was concentrated in vacuo to give
 (3S)-3-amino-4-oxobutyric acid tert-butyl ester
- 20 semicarbazone. The 0 °C solution of the above residue and 603a (200 mg, 0.42mmol) in 5 ml of DMF and 5 ml of CH₂Cl₂ was treated with 1-hydroxybenzotriazole (57 mg, 0.42mmol) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (98 mg, 0.51 mmol).
- 25 The reaction was stirred at RT for 18 h, poured onto

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EtOAc (75 ml) and washed with aq. 0.3 N KHSO₄, sat. aq. NaHCO₃ and sat. aq. NaCl, dried over NaSO₄ and concentrated *in vacuo*. Chromatography (flash, SiO₂, 0% to 4% MeOH/0.1% NH₄OH/CH₂Cl₂) to give 240 mg (83%) of

<u>Step B.</u> 604a was stirred with 10 ml of 33% TFA/ H_2O for 4 h and concentrated *in vacuo*. The residue was dissolved in 7 ml of MeOH/acetic acid/37% aq, formaldehyde (5:1:1) and stirred for 18 h.

- 10 Chromatography (Reverse Phase C18, 4.4 π m ID x 25 cm, 15% to 70% CH₃CN/0.1% TFA/H₂O) gave 32 mg (16%) of **605a** as a white solid: 1 H NMR (CD₃OD, existing as diastereomers of the hemiacetal) δ 7.85-7.78 (2H, d), 7.5-7.32 (6H, m), 7.32-7.28 (1H, m), 7.18-6.98 (5H, m),
- 15 4.92-4.85 (2H, m), 4.5-4.32 (2H, m), 4.31-4.20 (2H, m), $3.7-3.6 \ (1H, m), \ 2.90-2.75 \ (2H, m), \ 2.65-2.5 \ (1H, m), \\ 2.48-2.25 \ (3H, m).$

The following compounds were prepared by a similar method:

- 20 (3s)-3-[(3s)-2-Oxo-3-benzoylamino-5-phenylacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino|4-oxo-butyric acid (605b). 148 mg (33%) as a white solid: ¹H NMR(CD₃OD) & 7.9-6.9 (m, 16H), 4.9 (s, 2H), 4.5 (m, 1H), 4.4 (m, 2H), 3.75 (s, 1H), 3.6 (dd, 1H), 3.45 (dd, 1H), 2.7 (m, 1H), 2.5 (m, 1H).
 - (3S)-3-[(3S)-2-0xo-3-benzoylamino-5-benzoyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]4-oxo-butyric acid (605c). 319 mg (56%) as a white solid:

 1H NMR(CD₂OD) & 7.9-6.9 (m, 16H), 5.1 (m, 1H), 4.9 (dd,

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1H), 4.7 (m, 1H), 4.6 (dd, 1H), 4.4 (m, 2H), 4.05 (m, 1H), 2.7 (m, 1H), 2.5 (m, 1H).

(3S) -3-[(3S) -2-Oxo-3-benzoylamino-5-acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]4-oxo-5 butyric acid (605d). 190 mg (38%) as a white solid:

1H NMR(CD3OD) & 1.9(d, H), 2.4(m, 1H), 2.65(m, 1H), 3.7(m, 1H), 4.25(m, 1H), 4.45(m, 2H), 4.8-5.05(m, 3H),

(3S)-3-[(3S)-2-0xo-3-benzoylamino-5-methoxyacetyl-

10 2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]4-oxo-butyric acid (605e). 250 mg (78%) 1 H NMR (CD₃OD) δ 1.87 (bs), 1.95 (s, 2H), 2.1 (bs), 2.4 (m, 2H), 2.65 (m, 2H), 3.59 (bs), 3.75 (bs), 3.87 (bs), 4.19 (m), 4.37 (m), 4.50-4.78 (bm), 4.92 (m), 5.27

15 (bs), 7.41-7.58 (m, 7H), and 7.87 ppm (d, 2H).

7.3-7.7(m, 7H), 7.9(d, 2H).

(3S) -3-[(3S) -2-0xo-3-benzoylamino-5-(3-methylbutyryl) -2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-

acetylamino]4-oxo-butyric acid (605f). 210.5 mg (46%) as a white solid: $^1{\rm H}$ NMR(CD₃OD) δ 7.9-7.4 (m, 9H), 5.1

20 (m, 1H), 4.9 (m, 1H), 4.6 (dd, 1H), 4.4 (m, 2H), 4.1 (d, 1H), 3.8 (m, 1H), 3.5 (q, 1H), 2.7 (m, 1H), 2.5 (m, 1H), 2.0 (m, 3H), 1.2 (t, 1H), 0.9 (d, 3H), 0.8 (d, 3H).

(3S) -3-[(3S) -2-Oxo-3-benzoylamino-5-acetoacetyl-

25 2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1acetylamino]4-oxo-butyric acid (605g). 81 mg (19°) as
a white solid: ¹H NMR(CD₃OD) δ 7.9-7.3 (m, 11H), 4.94.8 (m, 2H), 4.6-4.4 (m, 3H), 4.3 (m, 1H), 3.75 (g,

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1H), 3.55 (d, 1H), 2.7 (m, 1H), 2.5 (m, 1H), 2.05 (s, 3H).

- (3S)-3-[(3S)-2-0xo-3-benzoylamino-5-methyloxalyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-
- 5 acetylamino]4-oxo-butyric acid (605h). 227 mg (54%) of a white solid: ¹H NMR(CD₃OD) δ 2.5 (m, 1H), 2.7 (m, 1H), 3.55 (s, 3H), 3.8-4.0 (m, 2H), 4.4 (m, 1H), 4.6-4.8 (m, 2H), 4.95 (d, 1H), 5.1 (m, 1H), 7.3-7.7 (m, 7H), 7.9 (d, 2H), 8.6 (d, 1H).
- 10 (3S)-3-[(3S)-2-Oxo-3-benzoylamino-5-acetylcarbonyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]4-oxo-butyric acid (605i). 150 mg (37%) as a white solid:

 1H NMR(CD₃OD) δ 7.9-7.3 (m, 12H), 5.1 (m, 1H), 4.65 (t, 1H), 4.55 (dd, 1H), 4.35 (m, 1H), 4.1 (d, 1H), 3.9 (q, 1H), 3.45 (q, 1H), 2.7 (m, 1H), 2.5 (m, 1H), 2.25 (s, 3H).
- (3S)-3-[(3S)-2-0xo-3-benzoylamino-5-methoxycarbonyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]4-oxo-butyric acid (605j). 234 mg (44%) as 20 a white solid: ¹H NMR(CD₃OD) & 7,9-7,4 (m, 12H), 5.0 (m, 1H), 4.8-4.5 (m, 3H), 4.4 (m, 1H), 4.3 (t, 1H), 3.9-3.75 (m, 2H), 3.6 (s, 3H), 2.7 (m, 1H), 2.5 (m, 1H).
- (3s)-3-[(3s)-2-Oxo-3-benzoylamino-5-methanesulfonyl25 2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1acetylamino]4-oxo-butyric acid (605m). 64.5 mg (34%)
 as a white solid: HNMR (DMSO-d₆, exisiting as
 diastereomers of the hemiacetal 6 open form of the
 aldehyde) 8 9.48 (0.2H, s), 8.85-8.72 (1H, m), 8.65-

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8.60 (0.8 H, d), 8.30-8.26 (0.2 H, d), 7.95-7.88 (2H,d), 7.6-7.45 (6H, m), 7.44-7.38 (1H, m), 5.78-5.75 (0.2H, d), 5.48 (0.6H, s), 4.85-4.70 (2H, m), 4.62-4.54 (1H, d), 4.50-4.40 (2H, m), 4.25-4.14 (1H, m), 3.9-3.85 (1H, m), 3.16 (3H, s), 3.05-2.3 (2, m).

(3S)-3-[(3S)-2-Oxo-3-benzoylamino-5-(naphthlene-2-carbonyl)-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino|4-oxo-butyric acid (605n). 103 mg (17%) as a white solid: ¹H NMR (CD₃OD) & 1,9(s, 3H), 2.5(m, 1H), 2.65(m, 1H), 3.75(m, 1H), 4.3(m,1H), 4.5-4.7(m, 3H), 4.85-5.1(m, 2H), 7.3-7.65(m, 6H), 7.85-8.05(m, 4H), 8.45(s. 1H).

(3S) -3-[(3S) -2-0xo-3-benzoylamino-5-acetyl-2,3,4,5-tetrahydro-7,9-dimethyl-1H-1,5-benzodiazepin-1-

- 15 acetylamino)4-oxo-butyric acid (605o). 42 mg (12%) as a white solid: ¹H NMR (CD₃OD, existing as diastereomers of the hemiacetal) δ 7.85-7.74 (2H, m), 7.5-7.44 (1H, m), 7.43-7.35 (4H, m), 5.6-5.05 (2H, m), 4.82-4.42 (2H, m), 4.40-3.95 (2H, m), 3.6-3.5 (1H, m), 2.7-2.38 (2H, m), 2.32 (3H, s), 2.27 (3H, s), 1.92 (3H, s).
- (3s)-3-[(3s)-2-Oxo-3-benzoylamino-5-benzylaminocarbonyl-2,3,4,5-tetrahydro-1H-1,5-benzo diazepin-1-acetylamino]d-oxo-butyric acid (605p). 165
 25 mg (37%) as a white solid: ¹H NMR (CD₃OD) δ 2.45(m, 1H), 2.7(m, 1H), 3.8(m, 1H), 4.15-4.5(m, 4H), 4.5-4.75(m, 2H), 4.8-5.0(m, 2H), 7.1-7.7(m, 12H), 7.9(d, 2H).

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(3S)-3-[(3S)-2-Oxo-3-benzoylamino-5-[(3R,S) 3-tetrahydrofuranylmethyoxycarbonyl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]4-oxo-butyric acid (605q). 210 mg (66%) ¹H NMR (CD₃OD) δ 1.95 (s, 2H), 5 2.4 (m, 2H), 2.65 (m, 2H), 3.29 (s, 3H), 3.78 (m), 3.87 (bs), 4.0 (d, 1H), 4.32 (m), 4.50-4.15 (m), 4.95 (m),

(3s) -3-[(3s) -2-0xo-3-benzoylamino-5-(4-pyridylacetyl) -2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-

5.27 (bs), 7.45-7.65 (m, 7H), and 7.89 ppm (d, 2H).

10 acetylamino]4-oxo-butyric acid (605s). 128 mg (19%) as a white solid: ¹H NMR(CD₃OD) & 8.5-7.4 (m, 13H), 5.0 (m, 1H), 4.7 (m, 1H), 4.5 (m, 2H), 4.45-4.4 (m, 3H), 3.8-3.7 (m, 2H), 2.7 (m, 1H), 2.5 (m, 1H).

(3S)-3-[(3S)-2-0xo-3-benzoylamino-5-(3-

(35) 3-[(35) 2-0xo-3-benzylamino-5-(3-phenylpropionyl)-2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]4-oxo-butyric acid trifluoroacetic acid salt (605v). 88 mg (28%) as a white solid: ¹H NMR

25 (CD₃OD) δ 7.63-7.51 (2H, m), 7.5-7.35 (7H, m), 7.25-7.10 (3H,m), 7.1-7.02 (2H, m), 5.04-4.96 (1H, m), 4.75-4.57 (2H, m), 4.38-4.17 (2H,m), 4.24-4.12 (2H, m), 4.10-4.07 (1H, d), 4.88-4.7 (1H, m), 2.90-2.80 (2H, m), 2.78-2.63 (1H,m), 2.55-2.35 (2H, m), 2.34-2.22 (1H, m).

 $\label{eq:the_compounds} \mbox{ Table 13 are described}$ below.

Table 13

	Ħ	R ₂	R ₃	R ₄	R ₆	R ₇
ĺ	609a	Н	PhCH ₂ CH ₂ C(0)	PhCH ₂ CH ₂ C(O)	C1	Cl.
	609ь	Н	CH ₃ C(O)	PhC(O)	Cl	Cl

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(3s) -3-[(3s) -2-0xo-3-(3-phenylpropionylamino) -5-(3-phenylpropionyl) -2,3,4,5-tetrahydro-1H-1,5-benzodiazepin-1-acetylamino] -4-(5,7-dichlorobenzoxazol-2-yl) -4-oxo-butyric acid (609a).

5 Step A. A solution of 204 (223 mg, 0.5 mmol) and 603r (300mg; 0.36 mmol) in 4 ml of DMF and 4 ml of CH₂Cl₂ was treated with (Ph₃P)₂PdCl₂ (10 mg), 1-hydroxybenzotriazle (135 mg, 1.0 mmol) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (115 mg, 0.6 mmol). Tri-n-butyl tin hydride (219 mg, 0.75 mmol) was added dropwise to the reaction and stirred for 18 h. The reaction was poured onto EtoAc and washed with ag. 10% NaHSO₄, sat. ag. NaHCO₃ and sat. ag. NaCl, dried over Na₂SO₄ and concentrated in vacuo. Chromatography (flash, SiO₂, 0% to 50% EtoAc/hexane) gave 360 mg (86%) of 607a as a foam.

Step B. A solution of 607a (360 mg) in 5 ml of CH₂Cl₂
was added dropwise to a suspension of 1,1,1-triacetoxy1,1-dihydro-1,2-benziodioxol-3(1H)-one (362 mg, 0.85
20 mmol) in 20 ml of CH₂Cl₂. The reaction was stirred for
4.5 h, diluted with CH₂Cl₂ and washed with a 1:1
mixture of sat. aq. NaHCO₃/sat. aq. Na₂S₂O₃, sat. aq.
NaHCO₃ (2x) and sat. aq. NaCl, dried over Na₂SO₄ and
concentrated in vacuo. Chromatography (flash, SiO₂,
25 20% EtOAc/CH₂Cl₂) gave 340 mg (95%) of the ketone 608a.

Step C. 608a (300 mg, 0.36 mmol) was dissolved in 25 ml of 25% TFA/CH₂Cl₂ and stirred at RT for 5 h and concentrated in vacuo. Chromatography (flash, SiO₂, 0 to 5% MeOH/CH₂Cl₂) gave 118 mg (42%) of 609a as a white 30 solid: ¹H NMR (CD₃OD) & 7.62-6.65 (16H, m), 4.85-4.7

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(1H, m), 4.68-4.42 (2H, m), 4.40-4.15 (2H, m), 3.48-3.28 (1H, m), 3.0-2.9 (1H, m), 2.9-2.6 (4H, m), 2.55-2.18 (3H, m), 2.16-1.96 (2H, m).

(3S)-3-[(3S)-2-0xo-3-benzoylamino-5-acetyl-2,3,4,5-

5 tetrahydro-1H-1,5-benzodiazepin-1-acetylamino]-4-(5,7-dichlorobenzoxazol-2-yl)-4-oxo-butyric acid (609b) was prepared from 603d in a similar manner as 609a to give 287 mg (438 overall yield) as white solid: 1 H NMR (DMSO- 1 G) δ 1.6(s, 3H), 2.7-3.1(m, 2H), 3.45(m, 1H),

10 4.4(t, 1H), 4.7(m, 2H), 4.95(m, 1H), 5.2, 5.4(2s, 1H), 7.2-7.65(m, 8H), 7.9(d, 2H), 8.8(t, 1H), 8.9,9.1(2s, 1H), 12.6(br, 1H).

$$R_{2}$$
 R_{2}
 R_{3}
 R_{2}
 R_{4}
 R_{2}
 R_{4}
 R_{5}
 R_{5

612

(3S)-3-[(3S)-2-0xo-3-benzoylamino-5-methanesulfonyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]-5-(2,6-dichlorobenzoyloxy)-4-oxo-pentanoic acid (612) was prepared by a method similar as 607a

(Steps A and C only) using 603m (150 mg, 0.36 mmol)

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instead of **603r** and (3*S*)-3-(allyloxycarbonylaminc)-4oxo-5-(2,6-dichlorobenzoyl-oxy)pentanoic acid t-butyl
ester (110; 160 mg, 0.36 mmol, WO 93/16710) instead of **606a** to give **612** (56%) as a white solid: ¹H NMR

5 (CDCl₃) 7.85-7.10 (12H, m), 5.4-4.65 (4H, m), 4.6-4.15
(4H, m), 3.10-2.72 (5H, s & m).

Example 13

 $\mbox{ Compounds } \mbox{ }$

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Syntheses of 619-635.

Step A. Synthesis of 614. TentaGel S@ NH $_2$ resin (0.16 mmol/g, 10.0 g) was placed in a sintered glass funnel and washed with dimethylformamide (3 X 50 mL),

- 5 10% (v/v) diisopropylethylamine (DIEA) in dimethylformamide (2 X 50 mL) and finally with dimethylformamide (4 X 50 mL). Sufficient dimethylformamide was added to the resin to obtain a slurry followed by 400 (1.42 g, 2.4 mmol, prepared from
- 10 (3S) 3-(fluorenylmethyloxycarbonyl)-4-oxobutryic acid t-butyl ester according to A.M. Murphy et. al. <u>x. Am.</u> <u>Chem. Soc.</u>, 114, 3156-3157 (1992)), 1hydroxybenzotriazole hydrate (HOBT H₂O; 0.367 g, 2.4 mmol), O-benzotriazole-N,N,N,N'-tetramethyluronium
- 15 hexafluorophosphate (HBTU; 0.91 g, 2.4 mmol), and DIEA (0.55 mL, 3.2 mmol). The reaction mixture was agitated overnight at room temperature using a wrist arm shaker. The resin was isolated on a sintered glass funnel by suction filtration and washed with dimethylformamide (3
- 20 X 50 mL). Unreacted amine groups were then capped by reacting the resin with 20% (v/v) acetic anhydride/dimethylformamide (2 X 25 mL) directly in the funnel (10 min/wash). The resin was washed with dimethylformamide (3 X 50 mL) and dichloromethane (3 X
- 25 50 mL) prior to drying overnight in vacuo to yield 614 (11.0 g, quantitative yield).

Step B. Synthesis of 616. Resin 614 (3.0 g, 0.16 mmol/g, 0.48 mmol) was swelled in a sintered glass funnel by washing with dimethylformamide (3 X 15 mL). 30 The Fmoc protecting group was then cleaved with 25%

(v/v) piperidine/dimethylformamide (15 mL) for 19 min

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(intermittent stirring) and then for 20 min with fresh piperidine reagent (15 ml). The resin was then washed with dimethylformamide (3 X 15 ml), followed by N-methypyrrolidone (2 X 15 mL). After transferring the resin to a 100 mL flask, N-methypyrrolidone was added to obtain a slurry followed by 603u (0.736 g, 0.72 mmol), HOBT'H2O (0.112 g, 0.73 mmol), HBTU (0.27 g, 0.73 mmol) and DIEA (0.26 mL, 1.5 mmol). The reaction mixture was agitated overnight at room temperature using a wrist arm shaker. The resin work-up and capping with 20% (v/v) acetic anhydride in dimethylformamide were performed as described for 614 to yield 616 (3.13 g, quantitative yield).

Step C. Synthesis of 617. This compound was

prepared from resin 616 (0.24 g, 0.038 mmol) using an
Advanced ChemTech 396 Multiple Peptide synthesizer.

The automated cycles consisted of a resin wash with
dimethylformamide (3 X 1 mL), deprotection with 25%
(v/v) piperidine in dimethylformamide (1 mL) for 3 min

followed by fresh reagent (1 mL) for 10 min to yield
resin 617. The resin was washed with dimethylformamide
(3 X 1 mL) and N-methypyrrolidone (3 X 1 mL).

with a solution of 0.4M thiophene-3-carboxylic acid and 0.4M HOBT in N-methypyrrolidone (1 mL), a solution of 0.4M HBTU in N-methylpyrrolidone (0.5 mL) and a solution of 1.6M DIEA in N-methypyrrolidone (0.35 mL, and the reaction was shaken for 2 hr at room temperature. The acylation step was repeated.

30 Finally, the resin was washed with dimethylformamide (3 X 1 mL), dichloromethane (3 X 1 mL) and dried in vacuo.

Step D. Method 1. (624). Resin 617 was acvlated

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The aldehyde was cleaved from the resin and globally deprotected by treatment with 95% TFA/ 5% H2O (v/v, 1.5 mL) for 30 min at room temperature. After washing the resin with cleavage reagent (1 mL), the combined 5 filtrates were added to cold 1:1 ether:pentane (12 mL) and the resulting precipitate was isolated by centrifugation and decantation. The resulting pellet was dissolved in 10% acetonitrile/90% H2O/O.1% TFA (15 mL) and lyophilized to obtain crude 624 as a white 10 powder. The compound was purified by semi-prep RP-HPLC with a Rainin Microsorb™ C18 column (5 u, 21.4 x 250 mm) eluting with a linear acetonitrile gradient (5% - 45%) containing 0.1% TFA (v/v) over 45 min at 12

mL/min. Fractions containing the desired product were 15 pooled and lyophilized to provide 624 (10.0 mg, 54%).

Step D. Method 1A. Synthesis of 627. Following a similar procedure as method 1, resin 617 was acylated with 4-(1-fluorenylmethoxycarbonylamino)benzoic acid and repeated. The Fmoc group was removed as described in Step C and the free amine was acetylated with 20 (V/V) acetic anhydride in dimethylformamide (1 mL) and 1.6M DIEA in N-methylpyrrolidone (0.35 mL) for 2 hr at room temperature. The acetylation step was repeated. Cleavage of the aldehyde from the resin gave 627 (4.2

Step D. Method 2. Synthesis of 632. Following a similar procedure as method 1, resin 617 was acylated with 0.5M cinnamoyl chloride in N-methypyrrolidone (1 mL) and 1.6M DIEA in N-methypyrrolidone (0.35 mL) for 2 may be a room temperature. The acylation step was

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repeated. Cleavage of the aldehyde from the resin gave $632 \ (11.1 \ \text{mg, } 58\%)$.

Step D. Method 3. Synthesis of 629. Following a similar procedure as method 1, resin 617 was reacted 5 with 1.0M benzenesulfonyl chloride in dichloromethane (0.5 mL) and 1M pyridine in dichloromethane (0.60 mL) for 4 hr at room temperature. The reaction was repeated. Cleavage of the aldehyde from the resin 629 (4.7 mg, 24%).

10 Analytical HPLC methods:

(1) Waters DeltaPak C18, 300A (5u, 3.9 X 150 mm). Linear acetonitrile gradient (5% - 45%) containing 0.1% TFA (v/v) over 14 min at 1 mL/min.

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Syn. Method	1	1	8		
MS Syn. (M+H) + Method	532	532	(M+Na) +		
HPLC RT min	11.71 (1)	531.53 10.44 (1)	11.57 (1) (M+Na)+		
MM	531.53	530.54			
MF	C27H25N5O7	C27H25N507	C28H26N4O7		
Structure	O N N N N N N N N N N N N N N N N N N N	N N N N N N N N N N N N N N N N N N N			
Ompd.	619	620	621		

MS Syn. (M+H)+ Method	H	ed.	, -1	
MS (M+H) +	(M+Na) + 569	(M-) 716	487	
HPLC RT min	10.19 (1) (M+Na)+ 98% 569	15.8 (1)	8.39 (1) 98 ₃	
MM	546.54	486.51		
MF	C28H26N4O8	C39H3ZN4O10	C22H22N407S	
Structure			STATE THE PART OF	
Cmpd.	622	623	624	

Cmpd.	Structure	Σ	MIN	HPLC RT	SW	Syn.
				min	+ (H+W)	(M+H) + Method
625	HO NO PER	C23H25N5O7S	515.55	7.60 (1)	51.6	good
626	HOHE OF THE PART O	C25H26N4O8	510.51	7.58 (1)	511	
627		C26H27N5O8	537.53	7.96 (1)	538	ΔI

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Cmpd.	Structure	MF	ММ	HPLC RT min	MS (M+H) +	MS Syn. (M+H)+ Method
628	H N N N N N N N N N N N N N N N N N N N	C25H24N409	524.49	9.50 (1)	525	
629	H	C23H24N408S	516.53	9.85 (1) 98%	517	m
630		C25H26N4O7	494.51	9.25 (1) 98§	495	7
631	ST S	C24H26N4O8S	530.56	10.19 (1)	531	м

Cmpd.	Structure	Æ	ММ	HPLC RT	MS	Syn.
				min	+ (H+W)	(M+H) + Method
632		C26H26N4O7	506.52	10.99 (1) 98%	507	2
633		C25II26N408	510.51	11.48 (1) 988	511	2
634	O O O O O O O O O O O O O O O O O O O	C22H26N409	490.47	6.87 (1) 988	491	5
635	HO W H	C25H24N408	508.49	10.03 (1) 98%	509	1

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Example 14

Compounds 1605a-j, 1605m, 1605n, 1605p, 1605t, and 1605v were synthesized as described below.

(3S) N-(2-Oxo-3-tert-butoxycarbonylamino-2,3,4,5-5 tetrahydro-1H-pyrido [3,4-b][1,4-diazepine (1600).

Step A. (2S) 2-tert-Butoxycarbonylamino-3-(3-nitropyridin-2-ylamino)propionic acid was prepared by a similar method as (2S) 2-tert-butoxycarbonylamino-3-(2-nitrophenyl-amino)propionic acid in Step A of the synthesis of 600a/103, except that 3-chlorc-3-nitro pyridine was used instead of 2-

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fluoronitrobenzene, to give 4.05 g (64%) of a yellow solid.

Step B. (2S) 2-tert-Butoxycarbonylamino-3-(3-aminopyridin-2-ylamino)propionic acid was prepared by

5 a similar method to (2S) 2-tert-Butoxycarbonylamino-3-(2-aminophenylamino)-propionic acid in Step B of the synthesis of 600a/103 to give 3.68 g (quant.) as a dark solid.

Step C. (2S) 2-tert-Butoxycarbonylamino-3-(3-

- aminopyridin-2-ylamino)propionic acid methyl ester. A
 solution of (2S) 2-tert-Butoxycarbonylamino-3-(3aminopyridin-2-ylamino)-propionic acid (360 mg, 1.21
 mmol) and MeOH (59 mg, 1.82 mmol) in anhydrous CH₂Cl₂
 (20 ml) was treated with 4-dimethylaminopyridine
- 15 (DMAP, 163 mg, 1.33 mmol) and 1-(3dimethylaminopropyl)-3-ethylcarbodiimide
 hydrochloride (280 mg, 1.45 mmol). The reaction was
 stirred for 18 h, diluted with EtOAc (150ml), washed
 with water (2x), sat. aq. NaHCO3, and sat. aq. NaCl,
- 20 dried over Na_2SO_4 and concentrated in vacuo. Chromatography (flash, SiO_2 , 0 to 5% MeOH/CH $_2Cl_2$) gave 250 mg (67%) of the title compound as a light tan solid.

Step D. (3S) N-(2-Oxo-3-tert-butoxycarbonylamino-

- 25 2,3,4,5-tetrahydro-1H- pyrido[3,4-b][1,4-diazepine (1600). A solution of (2S) 2-tert-butoxycarbonylamino-3-(3-aminopyridin-2-ylamino)prop ionic acid methyl ester (70 mg, 0.225 mol) and 25-sodium methoxide/MeOH (130 pl, 0.56 mmol) in
- 30 anhydrous MeOH (4 ml) was heated at 60°C for 16 h.

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The reaction was concentrated in vacuo, the residue dissolved in 2 ml of $\rm H_2O$ and extracted with EtOAc (3x). The combined extracts were dried over $\rm Na_2SO_4$ and concentrated in vacuo. Chromatography (flash, 5 $\rm SiO_2$, 0 to 3% MeOH/CH₂Cl₂) gave 7.5 mg (3%) of 1600 as a light tan solid: $^1\rm H$ NMR (CD₃OD) δ 7.96-7.92 (1H, d), 7.75-7.65 (1H, br. s), 7.14-7.08 (1H, d), 6.73-6.65 (1H, m), 5.83-5.75 (1H, br. s), 5.4-5.25 (1H, br. s), 4.6-4.5 (1H,m), 3.95-3.84 (1H, m), 3.55-10 3.48 (1H, m), 1.4 (9H, s)

Step E. 1601 is prepared from 1600 following the method in Step D for the preparation 600a/103.

Synthesis of 1603. 1603 is prepared from 1601 following the methods for the synthesis of 603 from 15 600.

PCT/US96/20843 WO 97/22619

Synthesis of 1605. 1605 is prepared from 1603 by methods described for the synthesis of 605 from 603.

Table 15

	1605	R ₃	R ₄
5	а	PhCH ₂ CH ₂ CO	PhCO
	ь	PhCH ₂ CO	PhCO
	c	PhCO	PhCO
	d	CH3CO	PhCO
	е	CH3OCH2CO	PhCO
10	f	(CH ₃) ₂ CHCH ₂ CO	PhCO
	g	CH₃COCH₂CO	PhCO
	h	сн ₃ ососо	PhCO
	i	CH3COCO	PhCO
	J	СН30СО	Ph.CO
15	m	CH3SO3	PhCO
	n	CH ₃ CO	Naphthyl-2-CO
	p	PhCH2NHCO	PhCO

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t	3-CH ₃ PhCH ₂ CO	PhCO
v	PhCH ₂ CH ₂ CO	PhCH ₂

Example 15

Compounds 1610-1621 are prepared from 1600

5 by methods similar to the methods used to prepare compounds 619-635 from 600a/103 and 600b.

- 518 -

wherein for compounds 1610-1621,

a
$$R_3 = CH_3C(O) - b$$
 $R_3 = CH_3OCH_2C(O) - :$

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Example 16

Compounds comprising scaffolds (ell), (yl), (y2), (z), and (el2) may be synthesized as described below.

5 Synthesis of Scaffold R_1 , wherein R_1 is (e11) and wherein Y_2 is =0.

- 520 -

Synthesis of Scaffold $R_1,$ wherein R_1 is (y1) and wherein Y_2 is =0.

- 521 -

Synthesis of Scaffold $R_1,$ wherein R_1 is (y2) and wherein Y_2 is H_2 and X_7 is O.

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Synthesis of Scaffold $R_1,$ wherein R_1 is $(\gamma 2)$ and wherein Y_2 is =0 and X_7 is NH.

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Synthesis of Scaffold $R_1\,,$ wherein R_1 is (y2) and wherein Y_2 is H_2 and X_7 is NH.

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Synthesis of Scaffold \mathbf{R}_1 , wherein \mathbf{R}_1 is (z) and wherein \mathbf{Y}_2 is O.

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Synthesis of Scaffold $R_{1}\,,$ wherein R_{1} is (e12) and wherein Y_{2} is =0.

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Example 17

The preparation of compounds 2001, 2002, 2100a-e, and 2201 is described below.

(15,95) 9-Benzovlformylamino-6,10-dioxo-

- 5 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a]-[1,2] diazepine-1-carboxylic acid (2000). To a solution of t-butyl 9-amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxylate (GB 2,128,984; 340 mg, 1.15 mmol) in
- 10 CH₂Cl₂ was added benzoylformic acid (260 mg, 1.7 mmol), HOBT (230 mg, 1.7 mmol) and EDC (340 mg, 1.7 mmol). The resulting mixture was stirred at ambient temperature for 16 hours, poured into IN HCl and extracted with CH₂Cl₂. The organic extracts were

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further washed with saturated NaHCO $_3$, dried over MgSO $_4$ and concentrated to afford 1999 as a pale yellow solid. The solid was dissolved in CH $_2$ Cl $_2$ (25 ml) and TFA (25 ml) and stirred overnight and 5 concentrated in vacuo to give 560 mg of 2000 as an

[1S,9S(2RS,3S)] 9-Benzoylformylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-N-(2(R,S)-benzyloxy-5oxotetrahydrofuran-3-yl)-6H-pyridazino[1,2-a][1,2]10 diazepine-1-carboxamide (2001), was synthesized from
2000 by methods similar to compound 213e to afford
410 mg (63%) of 2001 as a white solid: ¹H NMR
(CDCl₃; mixture of diastereomers) δ 8.25 (1H, d),
8.23 (1H, d), 7.78 (1H, dd), 7.65 (1H, bm), 7.50 (2H,
15 m), 7.40-7.25 (4H, m), 6.55 (1H, d), 5.57 (1H, d),
5.10 (1H, t), 5.05-4.95 (2H, m), 4.90, (1H, d), 4.80
(1H, d), 4.72 (1H, bm), 4.65 (1H, m), 4.55 (1H, m),
4.45 (1H, t), 3.25 (1H, m), 3.15 (1H, m), 3.00 (2H,
bm), 2,90 (1H, dd), 2.70 (1H, m), 2.47 (1H, dd), 2.45

 $\label{eq:continuous} [3S(1S,9S)] $$ 3-(9-Benzoylformylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamido)-4-oxobutanoic acid (2002).$

25 Compound 2001 (58.6 mg, 0.10 mmol) was treated with 15 ml of TFA/MeCN/water (1:2:3) and stirred at room temperature for 6.5 h. The reaction was extracted with ether. The aqueous layer was concentrated with azeotropic removal of the water using MeCN. The

20 (1H, m), 2.35 (1H, m), 2.00-1.75 (4H, m), 1.60 (1H,

bm).

30 product was suspended in CH_2Cl_2 , concentrated in vacuo and precipitated with ether to give 46.8 mg

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(99%) of **2002** as a white solid: ^{1}H NMR (CD₃OD) δ 9.05 (0.25H, d), 8.15 (1H, d), 7.68 (1H, t), 7.64 (0.25H, d), 7.55 (3H, t), 7.35 (0.5H, m), 5.22 (1H, t), 4.90 (1H, m), 4.58 (1H, dd), 4.50 (1H, m), 4.28 (1H, bm), 3.45 (1H, bm), 3.10 (1H, bt), 2.68 (1H, ddd), 2.60-2.45 (2H, m), 2.30 (1H, dd), 2.15-2.05 (2H, m), 1.90 (2H, bm), 1.68 (1H, bm).

[1s,9s(2Rs,3s)] 9-Benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-N-(2-isopropoxy-5-oxo10 tetrahydro-furan-3-yl)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide (2100a). A
solution of 214e (101 mg, 0.23 mmol) in isopropanol
(10 ml) was stirred at room temperature with a
catalytic amount of p-toluenesulfonic acid (10 mg).
15 After 75 minutes, the reaction mixture was poured
into saturated NaHCO3 and extracted with CH2Cl2. The
combined extracts were dried over NaySO4 and

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concentrated. Flash chromatography ($\rm SiO_2$, $\rm CH_2Cl_2$ to EtOAc) afforded 56 mg ($\rm S18$) of **2100a** as a white solid: $^{1}\rm H$ NMR ($\rm CDCl_3$; mixture of diastereomers) δ 7.9-7.8 (2H, m), 7.6-7.5 (1H, m), 7.5-7.4 (2H, m), 7.1 5 (0.5H, d), 6.9 (0.5H, d), 6.4 (0.5H,d), 5.6 (0.5H, d), 5.3 (0,5H, s), 5.2-5.1 (1H, m), 4.95 (0.5H, m), 4.75-4.5 (1.5H, m), 4.35 (0.5H, t), 4.1 (0.5H, m), 3.98 (0.5H, m), 3.3-2.75 (4H, m), 2.5-2.4 (2H, m), 2.25 (1H, m), 2.1-1.9 (3H,m) 1.75-1.55 (2H,m).

- 10 [3S(1S,9S)] 3-(9-Benzoylformylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4,4-diethoxy-butyric acid,
 ethyl ester (2100b). A solution of 214e (16 mg,
 0.036 mmol) in ethanol (2 ml) was stirred at room
- 15 temperature with a catalytic amount of ptoluenesulfonic acid (2 mg). After 5 days, the reaction mixture was poured into saturated NaHCO₃ and extracted with CH₂Cl₂. The combined extracts were dried over Na₂SO₄ and concentrated. Flash
- 20 chromatography (SiO₂, CH₂Cl₂:EtOAc 95:5 v/v) afforded 16 mg (81%) of 2100b as a white solid: ¹H NMR (CDCl₃) d 7.85-7.74 (2H,m), 7.55-7.38 (3H,m), 7.04-6.95 (1H,d), 6.61-6.48 (1H,d), 5.15-5.08 (1H,m), 4.63-4.53 (1H,m), 4.52-4.45 (1H,m), 4.42-4.35 (1H,m),
- 25 4.15-4.05 (2H,m), 3.74-3.60 (2H,m), 3.57-3.42 (2H,m), 3.39-3.26 (1H,m), 3.03-2.93 (1H,m), 2.92-2.82 (1H,m, 2.65-2.52 (2H,m), 2.42-2.25 (1H,m), 2.20-1.88 (4H,m), 1.76-1.50 (2H,m), 1.35-1.10 (9H,m).

[3S(1S,9S)] 3-(9-Benzoylformylamino-6,10-dioxo-

30 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4,4-dimethoxy-butyric acid

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methyl ester (2100c). A solution of 214e (165 mg, 0.37 mmol) in methanol (5 ml) was stirred at room temperature with a catalytic amount of ptoluenesulfonic acid (17.5 mg). After 4 days, the 5 reaction mixture was diluted with EtOAc and washed with 10% NaHCO3 (3x) and brine. The combined extracts were dried over Na2SO4 and concentrated. Flash chromatography (SiO2, EtOAc) afforded 127 mg (68%) of 2100c as a white solid: 1 H NMR (CDCl₃) δ 10 7.82 (2H, d), 7.55-7.50 (1H, m), 7.47-7.43 (2H, m), 7.02 (1H, d), 6.53 (1H, d), 5.20-5.10 (1H, m), 4.56-4.50 (1H, m), 4.45-4.50 (1H each, two m), 3.69 (3H, s), 3.41 (3H, s), 3.43 (3H, s), 3.35-3.25 (1H, m), 3.06-2.98 (1H, m), 2.94-2.83 (1H, m), 2.65-2.53 (2H, 15 m), 2.35-2.32 (1H, m), 2.15-2.07 (1H, m), 2.00-1.89 (3H, m), 1.75-1.56 (2H, m).

[3s(1s,9s)] 3-(9-Benzoylformylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamido)-4,4-diisopropoxy-butyric

- 20 acid, isopropyl ester (2100d). A solution of 214e (53 mg, 0.12 mmol) in isopropanol (5 ml) was stirred at 50 °C with a catalytic amount of p-toluenesulfonic acid (5 mg). After 3 days the reaction mixture was poured into saturated NaHCO $_3$ and extracted with
- 25 CH_2Cl_2 . The combined extracts were dried over Na_2SO_4 and concentrated. Flash chromatography (SiO₂, CH_2Cl_2 :EtOAc (4:1 to 1:1 v/v)) afforded 49 mg (68%) of 2100d as a white solid: 1H NMR (CDCl₃) δ 7.85 (2H, d), 7.50-7.43 (1H, m), 7.41-7.35 (2H, m), 7.02
- 30 (1H, d), 6.47 (1H, d), 5.13-5.07 (1H, m) 5.00-4.9 (1H, m), 4.61-4.55 (2H, m), 4.37-4.30 (1H, m), 3.80-3.70 (1H, m), 3.90-3.80 (1H, m), 3.42-3.35 (1H, m),

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3.03-2.93 (1H, m), 2.91-2.81 (1H, m), 2.62-2.50 (2H, m), 2.38-2.33 (1H, m), 2.12-2.06 (1H, m), 1.97-1.81 (3H, m), 1.70-1.60 (2H, m), 1.28-1.05 (18H, m).

2100e

[15,95(2R5,35)] 9-Benzovlamino-6,10-dioxo-

- 5 1,2,3,4,7,8,9,10-octahydro-N-(2-ethoxy-5-oxo-tetrahydro-furan-3-yl)-6H-pyridazino[1,2-a][1,2]-diazepine-1-carboxamide (2100e), was synthesized from 302 via methods used to synthesize 304a to afford 2100e, except ethanol and triethylorthoformate were
- 10 used instead of methanol and trimethylorthoformate. Chromatography (SiO₂, 5% ethanol/CH₂Cl₂) afforded 92 mg (68%) of a white solid: ¹H NMR (CDCl₃; mixture of diastereomers) δ 7.90-7.80 (2H, m), 7.60-7.50 (1H, m), 7.50-7.40 (2H, m), 7.30 (0.5H, d), 7.00 (0.5H, d)
- 15 d), 6.50 (0.5H, d), 5.50 (0.5H, d), 5.20-5.10 (1.5H, m), 4.95 (0.5H, m), 4.75-4.65 (0.5H, m), 4.65-4.50 (1H, m), 4.38 (0.05H, t), 4.00-3.90 (0.5H, m), 3.85-3.75 (0.5H, m), 3.75-3.65 (0.5H, m), 3.65-3.55 (6.5H, m), 3.30-2.70 (4H, m), 2.50-2.35 (2H, m), 2.30 (1H, m), 2.50-2.35 (2H, m), 2.50 (2H, m), 2.50-2.35 (
- 20 d), 2.15-1.90 (3H, m), 1.80-1.60 (2H, m), 1.25-1.20 (3H, two t)

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(3S)-3-[(3S)-2-oxo-3-(1-naphthoy1) amino-5-methoxyacety1-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (2201) was synthesized from 600b by the methods used 5 to synthesize 605b to afford 2201: ¹H NMR (CDCl₃) δ 8.30-8.22 (1H,m), 8.05-7.98 (1H, d), 7.96-7.93 (1H,m), 7.77-7.68 (1H,m), 7.67-7.40 (7H,m), 5.12-5.02 (1H,m), 4.98-4.41 (5H,m), 4.38-4.24 (1H,m), 4.07-4.00 (1H,d), 3.92-3.80 (2H,m), 3.32 (3H,s), 2.75-2.60 10 (1H,m), 2.58-2.35 (1H,m).

- 533 -Example 18

We obtained the following data for selected compounds of this invention using the methods described herein (Table 16, see Example 7; Tables 17 and 18, see

5 Examples 1-4). The structures and preparations of compounds of this invention are described in Examples 28-31.

Table 16 Comparison of Prodrugs for Efficacy in
LPS Challenged Mice: Inhibition of IL-18 Production.

10 The percent inhibition of IL-1β production after treatment with a compound of the invention is shown as a function of time after LPS challenge ("-" indicates that no value was obtained at that relative time).

15

Time of Compound Administration (relative to time of LPS challenge, PO, 50 mg/kg)

10	ITCIACIAE	CO CIME OF		Charlenge,	FO, 5	υι
	Compound	-2h	-1h	0h	+1h	7
	213f	(-4)	-	8	-	7
	213h	9	-	53	-	1
	213i	(-11)	-	62	-	i
20	213k	0	-	68	-	
	2131	(-18)	-	80	-	
	213m	26	-	42	-	
	2130	4	-	8	-	
	213p	21	-	29	-	
25	213q	17	-	91	-	ì
	213r	59	-	37	-	
	213x	0	-	78	-	,
	213y	29	-	50	-	
	214e	39	-	70	75	
		43	44	48	11	
			-	_=	47	
30	214k	12	-	31	-	
	2141	0	-	54	-	
	214m	0		17	-	
	214w	11	-	91	-	
	2641	0	-	23	-	
35	404	-	-	-	56	
		55		6	-	

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	Compound	-2h	-1h	0h	+1h
	412	0	-	0	-
		11	-	37	
	418	-	-	-	64
		25	-	52	
	434	-	-	-	80
		0		63	
_	450	0		35	
5	452	- 28	-	-	70
	456	-	· -	89	
	450	41	-	69	56 -
	470	0		36	
	471	0	_	34	
	475	0		15	-
10	481	27	-	0	-
	486	19	-	15	
	487	17	_	20	-
	528	25		67	-
	550f	0	-	50	-
15	550h	55		73	-
	550i	(-10)		23	
	550k	36	-	34	-
	5501	9	-	38	
	550m	45	-	52	-
20	550n	19	-	65	-
	550o	19	-	64	-
	550p	30	-	60	
	655	0		68	_
	656	31	-	16	
25	662	41	-	75	
	668	-	-	-	53
	695a	49	-	78	-
	1015	15	-	28	
	2001	64	62	58	55
30	2001a	10	-	16	
	2002	5	-	87	
	2100h	34	-	32	
	2100i	19	-	74	
	2100j	4.8	41	0	33
35	2100k	30	50	32	- 72
	21001	52		28	
	2100m	40		42	
	2100n	21	9	64	73
	21000	31	44	68	64

<u>Table 17</u> Data for selected compounds of this invention obtained using the methods described in Examples 1-4.

	Compound	UV- Visible Ki (nM)	Cell PBMC avg. IC50 (nM)			Clearance Rat, i.v.
	213f			3000		
5	213g			2200		
	213h			1500		
	213i			1100		
	213j					
	213k			2000		
10	2131			2000		
	213m			2500	i	
	2130		5000	3300		
	213p			<300		
	213q			<300		
15	213r			<300		
	213v	0.5	1,100	1100	41	23
	213x		4500	2500	1	
	213y			930		
	214j	4.2	2500	6000		
20	214k	0.2	500	580		22
	2141	6	1900	1100		12
	214m	1.5	530	2200 ;		33.4
	214w	0.6	620	370		15
	246b	30000	>30000		87	
25	2641			3000		
	265a	2600	25000			
	265c	1100	4500			32
	265d	500	1500			35
	265f	1200				24
30	280b		13000			
	280c	-	10000			86
	280d		25000			
	283b		1750			41
	283c		4000			50

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	Compound	UV- Visible Ki (nM)	Cell PBMC avg. IC50 (nM)	blood	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
	283d		>8000	10000		
	308c	3000				
	308d	3000				
	500	25	1800	1800		
5	501	2.5	1800	1600		
	505c		1500			
	505d		>20000			
	505f		550			
	510a	65	200		267	
10	510d	2300	>20000		i	
	511c	730	>20000		78	40
	528			2200		
	550f			1100	Ī	
	550h			1800		
15	550i	_		1400		
	550k			3000		
	5501			750		
	550m			2000		
	550n			<300		
20	550o		450	3000		
	550p			2900	1	
	550q			700		
	640	155	2250	3900	1	
	642	35	8000	2900		
25	645	150				
	650	550	4000	i		
	653	30	2300	6000		
	655					
	656	0.6	2100	1600		2.9
30	662	0.5	1800	800		2.75
	668	9	5200	3700		29
	669	14		10000		
	670			4500		
	671	5	2000	2500		33.2
35	677		,	610 :		-
	678	5	2700	2200		
	680					

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	Compound		Cell PBMC avg. IC50 (nM)	prood	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
	681	9	3000	5000		
	682			1300		
	683	400	>20000	>20000		
	684	15	5000	2800		
5	686	4	4000	9000		
	688a			3000		
	688b			1300		
	689a	0.8	910	2500		
	689b	2.2	600	2000		
10	690a			1600		
	690b					
	691a	2.1	2900	1200		9.9
	691b	11.5	1,900	1400		
	692a					
15	692b		-	1800		
	693		1			
	694	3	2600	2100		
	695a					,
	695b					
20	695c			2500		
	696	4.5	2000	2900		13
	700	275				
	701	90			1	
	702	45	>5000	20000		
25	703	5	1400	20000 .	1	
	704	30	2600	9800		
	705	5	2300	3200		
	706	5	2400	5800		
	707	180				
30	708	140				
	709	10	2100	14000		
	710	110				
	711	175	-			
	910	10	3400	3800		
35	911	9	3500	1900		
	912	10	4200	3800		_
	913	4.5	2400	7000		

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	Compound	UV- Visible Ki (nM)	Cell PBMC avg. IC50 (nM)		Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
	914	5.2	2600	2800		
	915	11.5	>8000	1900		
	918	7		1150		
	919	4	2000	4300		
5	920	16	2100	3000		
	921	8.5	1800	3000		
	1018	170	4000	5500		9.1
	1052	100	2500			16
	1053	27	2000	>20000		34
10	1056	170				17
	1075	120	5000	5500		14.5
	1095	360	6000			28
	1105	250	3500	3000		
	1106	75	4000	1700		
15	1107	65				
	1108	22	1400	2600		
	1109	80				
	1110	45				
	1111	18	6050	4400		
20	1112	3.5	1800	2300		
	1113	290				
	1114	125				
	1115	250				
	1116	215				
25	1117	35	1700	1300		
	1118	380				
	1119	515				
	1120	95			ŀ	
	1121	170				
30	1122	400				
į	1123	30	2,400	4500		
	1124	270			1	
1	1125	55	2300	9000		
1	2001a			3000		
35	2100f					
	2100g					
	2100h			2000		

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Compound	UV- Visible Ki (nM)	Cell PBMC avg. IC50 (nM)	human blood	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
2100i					
2100j	30000		12000		
2100k	520	4000	600		
21001		750	2200		
2100m					
2100n	670	770	4000		
2100o	670	1150	1500		

We obtained the following data for selected compounds of this invention (Table 18) using the 10 methods described herein (see Examples 1-4). The structures and preparations of compounds of this invention are described in Examples 28-31.

Table 18

5

	Cmpd.	Fluorescent Assay kinact m ⁻¹ s ⁻¹	Cell PBMC avg. IC50 (nM)	Whole human blood IC50 (nM)	Mouse, i.v.	Clearance Rat, i.v. ml/min/kg
15	286	370000	300	1600		119
	505 b	190000	1500	2100	161	196
	505 e	420000	9000	1000		

Example 19

In vivo acute assay for efficacy as 20 anti-inflammatory agent

Results in the Table 19 show that 412f, 412d and 696a inhibit LL-1 β production in LPS-challenged mice after oral adminstration using ethanol/PEG/water,

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 β -cyclodextrin, labrosol/water or cremophor/water as vehicles. The compound was dosed at time of LPS challenge. The protocol is described in Example 7.

Table 19 Inhibition (%) of IL-1 β production in LPS-5 challenged mice.

Compound	10 mg/kg	25 mg/kg	50 mg/kg
	dose	dose	dose
412f	17%	25%	32%
412e	5%	17%	61%
696a	0	45%	52%

10

Example 20 Mouse Carrageenan Peritoneal Inflammation

Inflammation was induced in mice with an

intraperitoneal (IP) injection of 10 mg carrageenan in
 0.5 ml of saline (Griswold et al., <u>Inflammation</u>, 13,
15 pp. 727-739 (1989)). Drugs are administered by oral
 gavage in ethanol/PEG/water, β-cyclodextrin,
 labrosol/water or cremophor/water vehicle. The mice are

- labrosol/water or cremophor/water vehicle. The mice are sacrificed at 4 hours post carrageenan administration, then injected IP with 2 ml of saline containing 5U/ml 20 heparin. After gentle massage of the peritoneum, a
- small incision is made, the contents collected and volume recorded. Samples are kept on ice until centrifuged (130 x g, 8 mins at 4 °C) to remove cellular material, and the resultant supernatant stored
- 25 at -20 °C. IL-1 β levels in the peritoneal fluid are determined by ELISA.

Results in the Table 20 show prodrug 412f inhibits ${\rm IL-1}\beta$ production in carrageenan-challenged mice after oral administration of drug. Compound 214e

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did not inhibit $IL-1\beta$ production when dosed orally at 50 mg/kg.

Table 20 Inhibition (%) of IL-1β production by 412f and 412d in carrageenan-challenged mice.

5	Dose	Compound 412f	Compound 412d
	(mg/kg)		1
	1	30%	0
	10	54%	32%
	25	49%	31%
10	50	73%	36%

75%

100

Example 21 Type II Collagen-induced Arthritis

53%

15 Type II collagen-induced arthritis was established in male DBA/1J mice at described Woolev and Geiger (Wooley, P.H., Methods in Enzymology, 162, pp. 361-373 (1988) and Geiger, T., Clinical and Experimental Rheumatology, 11, pp. 515-522 (1993)).

- 20 Chick sternum Type II collagen (4 mg/kg in 10 mM acetic acid) was emulsified with an equal volume of Freund's complete adjuvant (FCA) by repeated passages (400) between two 10 ml glass syringes with a gauge 16 double-hub needle. Mice were immunized by intradermal
- 25 injection (50 μ l; 100 μ l CII per mouse) of collagen emulsion 21 days later at the contra-lateral side of the tail base. Drugs were administered twice a day (10, 25 and 50 mg/kg) by oral gavage approximately 7 $\rm h$ apart. Vehicles used included ethanol/PEG/water, B-
- 30 cyclodextrin, labrosol/water or cremophor/water. Drug treatments were initiated within 2 h of the CII booster

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immunization. Inflammation was scored on a 1 to 4 scale of increasing severity on the two front paws and the scores are added to give the final score.

Results in the Figs. 12, 13 and 14 show 5 prodrugs 412f, 412d and 696a inhibit inflammation in collagen-induced arthritits in mice after oral adminstration. Compound 214e did not inhibit inflammation when dosed (50 mg/kg) once a day by oral gavage.

10 Example 22

In vivo bioavailability determination

The drugs (10-100 mg/kg) were dosed orally to rats (10 mL/kg) in ethanol/PEG/water, β -cyclodextrin, labrosol/water or cremophor/water. Blood samples were

- 15 drawn from the carotid artery at 0.25, 0.50, 1, 1.5, 2, 3, 4, 6, and 8 hours after dosing, centrifuged to plasma and stored at -70°C until analysis. Aldehyde concentrations were determined using an enzymatic assay. Pharmacokinetic analysis of data was performed
- 20 by non-linear regression using RStrip (MicroMath Software, UT). Drug availability values were determined as follows: (AUC of drug after oral prodrug dosing/AUC of drug after i.v. dosing of drug)x(dose i.v./dose p.o.) x100%.
- Results in Table 21 show that prodrugs 412f, 412d and 696a give significant blood levels of drug and have good drug availability when dosed orally. Blood levels of 214e were not detected when it was dosed orally.

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Compound	Dose	Cmax	Drug
	(mg/kg)	(µg/ml)	Availability (%)
412f	25	2.4	32
412d	25	2.6	35
696a	50	1.2	10
214e	45	0.2	0.9%

Example 23
ICE cleaves and activates pro-IGIF

10 ICE and ICE homolog expression plasmids

5

A 0.6 kb cDNA encoding full length murine pro-IGIF (H. Okamura et al., Nature, 378, p. 86 (1995) was ligated into the mammalian expression vector pCDLSRa (Y. Takebe et al., Mol. Cell Biol., 8, p. 466 (1988)).

Generally, plasmids (3 μg) encoding active ICE (above), or the three ICE-related enzymes TX, CPP32, and CMH-1 in the pCDLSRα expression vector (C. Faucheu et al., EMBQ, 14, p. 1914 (1995); Y. Gu et al.,

- 20 EMEO, 14, p. 1923 (1995); J. A. Lippke et al., J. Biol. Chem., 271, p. 1825 (1996)), were transfected into subconfluent monolayers of Cos cells in 35-mm dishes using the DEAE-dextran method (Y. Gu et al., EMBC J., 14, p. 1923 (1995)). Twenty-four hours later, cells
- 25 were lysed and the lysates subjected to SDS-PAGE and immunoblotting using an antiserum specific for IGIF (H. Okamura et al., Nature, 378, p. 88 (1995).
 Polymerase chain reaction was used to
 - introduce Nde I sites at the 5' and 3' ends of the
- 30 murine pro-IGIF cDNA using the following primers:
 GGAATTCCATATGGCTGCCATGTCAGAAGAC (forward) and
 GGTTAACCATATGCTAACTTTGATGTAAGTTAGTGAG (reverse). The

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resulting NdeI fragment was ligated into \underline{E} , \underline{coli} expression vector pET-15B(Novagen) at the NdeI site to create a plasmid that directs the synthesis of a polypeptide of 213 amino acids consisting of a 21-

- 5 residue peptide (MGSSHHHHHHSGG_VPRGSHM, where LVPRGS represents a thrombin cleavage site) fused in-frame to the N-terminus of pro-IGIF at Ala2, as confirmed by DNA sequencing of the plasmid and by N-terminal sequencing of the expressed proteins. E. coli strain BL21(DE3)
- 10 carrying the plasmid was induced with 0.8 mM isopropyl-1-thio-β-D-galactopyranoside for 1.5 hours at 37°C, harvested, and lysed by microfluidization (Microfluidic, Watertown, MA) in Buffer A (20 mM sodium phosphate, pH 7.0, 300 mM NaCl, 2 mM dithiothreitol,
- 15 10% glycerol, 1 mM phenylmethylsulfonyl fluoride, and 2.5 µg/ml leupeptin). Lysates were cleared by centrifugation at 100,000 x g for 30 min. (His) 6tagged pro-IGIF protein was then purified from the supernatant by Ni-NTA-agarose (Qiagen) chromatography 20 under conditions recommended by the manufacturer.

In Vitro pro-IGIF Cleavage Reactions

In vitro cleavage reactions (30 ul) contained 2 µg of purified pro-IGIF and various concentrations of the purified proteases in a buffer containing 20 mM 25 Hepes, pH 7.2, 0.1% Triton X-100, 2 mM DTT, 1 mM PMSF and 2.5 µg/ml leupeptin and were incubated for 1 hour at 37°C. Conditions for cleavage by granzyme 5 were as described previously (Y. Gu et al., J. Biol. Chem., 271, p. 10816 (1996)). Cleavage products were analyzed by SDS-PAGE on 16% gels and Coomassie Blue staining, and were subjected to N-terminal amino acid sequencing

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using an ABI automated peptide sequencer under conditions recommended by the manufacturer.

Kinetic Parameters of IGIF Cleavage by ICE

The kinetic parameters (kgat/KM, KM, and kgat)

- 5 for IGIF cleavage by ICE were determined as follows.

 35S-methionine-labeled pro-IGIF (3000 cpm, prepared by
 in vitro transcription and translation using, the TNT

 T7-coupled reticulocyte lysate system (Promega) and
 pro-IGIF cDNA in a pSP73 vector as template) were

 10 incubated in reaction mixtures of 60 ul containing 0.1
- to 1 nM recombinant ICE and 190 nM to 12 µM of unlabeled pro-IGIF for 8-10 min at 37°C. Cleavage product concentrations were determined by SDS-PAGE and PhosphoImager analyses. The kinetic parameters were
- 15 calculated by nonlinear regression fitting of the rate vs. concentration data to the Michaelis-Menten equation using the program Enzfitter (Biosoft).

IFN-v Induction Assavs

A.E7 Th1 cells (H. Quill and R. H. Schwartz,

20 J. Immunol., 138, p. 3704 (1987)) (1.3 x 10⁵ cells in
0.15 ml Click's medium supplemented with 10% FBs, 50 uM
2-mercaptoethanol and 50 units/ml IL-2) in 96-well
plates were treated with IGIF for 18-20 hours and the
culture supernatant were assayed for IFN-y by ELISA
25 (Endogen, Cambridge, MA).

Example 24

Processing of pro-IGIF by ICE in Cos Cells

Cos cells were transfected with various expression plasmid combinations as described in Example

30 23. Transfected Cos cells $(3.5 \times 10^5 \text{ cells in a } 35\text{-mm})$ dish) were labeled for 7 hours with 1 ml of methionine-

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free DMEM containing 2.5% normal DMEM, 1% dialyzed fetal bovine serum and 300 µCi/ml 35S-methionine (35S-Express Protein Labeling-Mix, New England Nuclear). Cell lysates (prepared in 20 mM Hepes, pH 7.2, 150 mM 5 NaCl, 0.1% Triton X-100, 5 mM N-ethylmaleimide, 1 mM PMSF, 2.5 µg/ml leupeptine) or conditioned medium were immunoprecipitated with an antiIGIF antibody that recognizes both the precursor and the mature forms of IGIF (H. Okamura et al., Nature, 378, p. 88 (1995)). 10 Immunoprecipitated proteins were analyzed by SDS-PAGE (polyacrylamide gel electrophoresis) and fluorography (Fig. 2A).

We also measured the presence of IFN-y inducing activity in the cell lysates and the 15 conditioned media of transfected cells (Fig. 2B). Transfected Cos cells $(3.5 \times 10^5 \text{ cells in a } 35\text{-mm dish})$ were grown in 1 ml medium for 18 hours. Media was harvested and used at 1:10 final dilution in the IFN-v induction assay (Example 23). Cos cell pellets from 20 the same transfection were lysed in 100 ul of 20 mM Hepes, pH 7.0, by freeze-thawing 3 times. Lysates were cleared by centrifugation as described above and were used at a 1:10 dilution in the assay.

Example 25

IGIF is a physiological substrate of ICE Wild type (ICE+/+) and ICE-/- mice were primed with heat-inactivated P. acnes, and Kupffer cells were isolated from these mice 7 days after priming and were then challenged with 1 µg/ml LPS for 30 3 hours. The amounts of IGIF in the conditioned media were measured by ELISA.

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Wild type or ICE-deficient mice were injected intraperitoneally with heat-killed p. acnes as described (H. Okamura et al., Infection and Immunity, 63, p. 3966 (1995)). Kupffer cells were prepared seven 5 days later according to Tsutsui et al. (H. Tsutsui et al., Hepato-Gastroenterol., 39, p. 553 (1992)) except a nycodenz gradient was used instead of metrizamide. For each experiment, Kupffer cells from 2-3 animals were pooled and cultured in RPMI 1640 supplemented with 10% 10 fetal calf serum and 1 µg/ml LPS. Cell lysates and conditioned medium were prepared 3 hours later.

Kupffer cells from wild type and ICE-/- mice were metabolically labeled with 35S-methionine as for Cos cells (described above in Example 24) except that 15 methionine-free RPMI 1640 was used in place of DMEM. IGIF immunoprecipitation experiments were performed on cell lysates and conditioned media and immunoprecipitates were analyzed by SDS-PAGE and fluorography as described in Example 23. See Fig. 3.

20

Example 26

Induction of IFN-v Production In Vivo LPS mixed with 0.5% carboxymethyl cellulose in PBS, pH 7.4, was administered to mice by intraperitoneal injection (30 mg/kg LPS) in a dose 25 volume of 10 ml/kg. Blood was collected every 3 h for 24 h from groups of three ICE-deficient or wild type mice. Serum IFN-y levels were determined by ELISA (Endogen).

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Example 27

IGIF and IFN-y Inhibition Assays

Inhibition of IGIF processing by ICE inhibitors was measured in ICE inhibition assays as 5 described herein (see Example 1 and Table 22).

Human PBMC Assays

Human buffy coat cells were obtained from

blood donors and peripheral blood mononuclear cells
(PBMC) were isolated by centrifugation in LeukoPrep
10 tubes (Becton-Dickinson, Lincoln Park, NJ). PBMC were
added (3 x 10⁶/well) to 24 well Corning tissue culture
plates and after 1 hr incubation at 37°C, non-adherent
cells were removed by gently washing. Adherent
mononuclear cells were stimulated with LPS (1 ug/ml)

15 with or without ICE inhibitor in 2 ml RPMI-1640-108 FBS. After 16-18 hr incubation at 37°C, IGIF and IFN-y were quantitated in culture supernatants by ELISA.

For example, we obtained the following data for compound 412 of this invention using the methods 20 described herein. The structure of compound 412 is shown below.

Table 22

compound	UV-Visible	Cell PBMC	
	K _i (nM)	avg. IC50 (nM)	
412	1.3	580	

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Example 28

Compounds of this invention may be prepared via various methods. The following illustrates a preferred method:

To a solution of A (1.1 equivalent) in CH₂Cl₂ (or DMF, or CH₂Cl₂:DMF (1:1)) is added triphenylphosphine (0-0.5 equivalent), a nucleophilic scavenger (2-50 equivalents) and tetrakistriphenylphosphine palladium(0) (0.05-0.1 equivalent) at ambient temperature under inert atmosphere (nitrogen or argon). After 10 minutes, the above reaction mixture is optionally concentrated, then a solution of acid A-I or A-II in CH₂Cl₂ (or DMF, or CH₂Cl₂:DMF (1:1)) is added followed by addition of HOBT (1.1 equivalent) and EDC (1.1 equivalent). The resulting reaction mixture is allowed to stir at ambient temperature 1 hour-48 hours to provide coupled products C-I or C-II.

Various nucleophilic scavengers may be used in the above process. Merzouk and Guibe, <u>Tetrahedron</u> 20 <u>Letters</u>, 33, pp. 477-480 (1992); Guihe and Balavoine, <u>Journal of Crganic Chemistry</u>, 52, pp. 4984-4993 (1987)). Preferred nucleophilic scavengers that may be used include: dimedone, morpholine, trimethylsilyl dimethylamine and dimethyl barbituric acid. More preferred nuclophilic scavengers are trimethylsilyl dimethylamine (2-5 equivalents) and dimethyl barbituric (5-50 equivalents). When the nucleophilic scavenger is trimethylsilyl dimethylamine, the above reaction mixture must be concentrated prior to addition of A-I or A-II.

Other compounds of this invention may be prepared by hydrolyzing compounds represented by C-I and C-II to compounds represented by H-I and H-II as described in the following scheme:

The hydrolysis may be carried out under various conditions, provided that the conditions include an acid and H₂O. Acids that may be used include p-toluensulfonic, methanesulfonic acid, sulfuric, perchloric, trifluoroacetic, and hydrochloric. For example, trifluoroacetic acid (1-90% by weight) or

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hydrochloric acid (0.1-30% by weight) in CH_3CN/H_2O (1-90% H_2O by weight) at between 0-50 °C may be used.

Example 29

Compounds 213f, 213g, 213h, 213i, 213j, 213k, 5 213l, 213m, 214f, 214g, 214h, 214i, 214j, 214k, 214l, 214m, 550f, 550g, 550h, 550i, 550j, 550k, 550l and 550m were prepared as follows.

[1S,9S(2RS,3S)]9-[(4-Dimethylaminobenzoyl)amino]-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213f),
was synthesized from 212f by the methods used to

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prepare 213e from 212e to afford 504 mg of 213f as a yellow solid, $^1{\rm H}$ NMR (CD₃OD) δ 1.10(br. m, 0.25H), 1.30(br. m, 2H), 1.50(br. m, 1H), 1.65(br. m, 1.5H), 1.80(br. m, 0.25H), 1.90(br. m, 0.25H), 1.95(br. m, 50.5H), 2.05(br. m, 0.25H), 2.15(m, 1H), 2.3(m, 1H), 2.5(br. m, 1H), 2.6(dd, 1H), 2.8(m, 1H), 3.1(br. s, 3H), 3.15(br. m, 1H), 3.32(br. s, 3H), 3.5(m, 1H), 4.5(br. m, 1H), 4.62(d, 0.25H), 4.72(m, 3H), 4.95(m, 1H), 5.1(br. t, 0.25H), 5.15(br. t, 0.75H), 5.7(d, 1H), 10 6.75(d, 2H), 7.35(br. s, 5H), 7.75(d, 2H).

[1S, 9S(2RS, 3S)]9-[(3-Dimethylaminobenzoyl) amino]-6, 10-dioxo-1, 2, 3, 4, 7, 8, 9, 10-octahydro-N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6H-

0.75H), 5.2(d, 1H), 6.95(d, 1H), 7.15(d, 1H), 7.25(br. 25 s, 1H), 7.3(br. t, 2H), 7.45(br. s, 6H).

[15,95(2R5,35)]9-[(3-Chloro-4-aminobenzoyl)amino]-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6H-

m, 2H), 4.95(br. m, 1H), 5.15(br. t, 0.25H), 5.2(br. t,

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213h),

30 was synthesized from 212h by the methods used to

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prepare 213e from 212e to afford 296 mg of 213h, $^1\mathrm{H}$ NMR (CDC1₃) δ 1.55-1.68 (m, 1H), 1.7-2.05 (m, 3H), 2.3-2.5 (m, 2H), 2.65-2.8 (m, 1H), 2.85-2.93 (m, 1H), 2.95-3.25 (m, 3H), 4.44-4.65 (m, 2H), 4.68-4.82 (m, 1H), 4.9-4.95 (d, 5 1H), 5.05-5.18 (m, 2H), 5.28 (s, 0.5H), 5.55-5.58 (d, 0.5H), 6.52-6.58 (d, 0.5H), 6.7-6.76 (m, 2H), 6.82-6.85 (d, 0.5H), 7.3-7.4 (m, 5H), 7.52-7.58 (m, 1H), 7.75 (s, 0.5H), 7.8 (s, 0.5H).

[1S, 9S(2RS, 3S)]9-[(4-Methoxybenzoyl)amino]-6,10-dioxo-

10 1,2,3,4,7,8,9,10-octahydro-N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213i),
was synthesized from 212i by the methods used to
prepare 213e from 212e to afford 1.1 g of 213i, 1/2 NMR

- 15 (CDC1₃) δ 1.55-2.05 (m, 6H), 2.26-2.5 (m, 2H), 2.68-2.82 (m, 1H), 2.85-2.92 (m, 1H), 2.95-3.25 (m, 2H), 3.82 (s, 1.5H), 3.85 (s, 1.5H), 4.4-4.65 (m, 2H), 4.78 (m, 1H), 4.88-4.95 (m, 1H), 5.05-5.23 (m, 1H), 5.28 (s, 0.5H), 5.55-5.58 (d, 0.5H), 6.6-6.65 (m, 1H),
- 20 6.8-6.84(m, 1H), 6.9-6.95(m, 3H), 7.3-7.45(m, 4H), 7.78-7.85(m, 2H).

[15,95(2R5,35)]9-[(3,5-Dichlorobenzoyl)amino]-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6H-

25 pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213j),
 was synthesized from 212j by the methods used to
 prepare 213e from 212e to afford 367 mg of 213j, ¹H NMR
 (CDCl₃) δ 1.55-2.05 (m, 12H), 2.25 (d, 1H), 2.35 (m, 1H),
 2.48 (m, 2H), 2.75 (m, 2H), 2.9 (m, 1H), 2.95-3.25 (π, 5H),
30 4.45 (t, 1H), 4.5-4.6 (m, 4H), 4.7 (m, 1H), 4.75 (d, 1H),

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4.88 (m, 1H), 5.05 (m, 2H), 5.15 (q, 1H), 5.3 (s, 1H), 5.58 (d, 1H), 6.5 (d, 1H), 6.9 (d, 1H), 7.05 (d, 1H), 7.25-7.35 (m, 5H), 7.6 (s, 2H), 7.7 (s, 2H).
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[1S,9S(2RS,3S)]9-[(3,5-Dichloro-4-

- 5 hydroxybenzoyl)amino]-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213k), was synthesized from 212k by the methods used to prepare 213e from 212e to afford 593 mg of 213k, ¹H NMR (CD₃OD) δ 1.5(m, 1H), 1.6-1.7(m, 2H), 1.75-1.95(m, 4H), 2.15(m, 2H), 2.3(m, 1H), 2.6(m, 1H), 2.7(m, 1H), 3.05(m, 2H), 3.15(m, 1H), 3.5(m, 2H), 4.45(m, 2H), 4.65(d, 1H), 4.7(m, 1H), 4.95(m, 1H), 5.15(m, 1H),
- 15 [1s,9s(2Rs,3s)]9=[(3-Chloro-4-acetamidobenzoy1)amino]6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-Benzyloxy-5oxotetrahydrofuran-3-y1)-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (2131),
 was synthesized from 2121 by the methods used to

5.4(s, 1H), 5.7(d, 1H), 7.3(m, 5H), 7.85(s, 2H).

- 220 prepare 213e from 212e to afford 133 mg of 2131, 1 H NMR (CDCl₃) δ 1.55-1.7 (m, 1H), 1.75-2.05 (m, 3H), 2.25 (s, 1.5H), 2.27 (s, 1.5H), 2.3-2.48 (m, 2H), 2.7-2.83 (m, 1H), 2.85-2.94 (dd, 1H), 2.95-3.25 (m, 2H), 4.42-4.65 (m, 2H), 4.66-4.85 (m, 1H), 4.89-4.95 (m, 1H), 5.05-5.18 (m, 2H).
- 25 5.32(s, 0.5H), 5.55-5.6(d, 0.5H), 6.48-6.55(d, 1H), 6.88-6.92(d, 1H), 7.0-7.04(d, 0.5H), 7.15-7.2(d, 0.5H), 7.3-7.4(m, 4H), 7.64-7.78(m, 2H), 7.88-7.94(m, 1H), 8.45-8.56(m, 1H).

[1S,9S(2RS,3S)]9-[(3.5-Dichloro-4-

30 methoxybenzovl)amino]-6,10-dioxo-1,2,3,4,7,8,9,10-

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octahydro-N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213m), was synthesized from 212m by the methods used to prepare 213e from 212e to afford 991 mg of 213m, ¹H NMR 5 (CDCl₃) δ 1.5-2.15 (m, 5H), 2.2-2.55 (m, 3H), 2.6-3.3 (m, 4H), 3.95(2s, 3H), 4.45-4.7 (m, 2H), 4.7-4.85 (m, 1H), 4.8504.95 (m, 1H), 5.05-5.25 (m, 1H), 5.3 (s, 0.5H), 5.6 (d, 0.5H), 6.55 (d, 0.5H), 7.75 (s, 1H), 7.85 (s, 1H).

- 10 [15,95(2RS,3S)]9-[(4-Dimethylaminobenzoyl)amino]-6,10dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-ethoxy-5oxotetrahydrofuran-3-yl)-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (550f),
- was synthesized from 212f by the methods used to
- 15 prepare 213e from 212e to afford 420 mg of 550f as an off white solid, ^{1}H NMR (CDCl $_{3}$) δ 1.2-1.25(br. τ , 3H), 1.35(m, 1H), 1.55(br. π , 1H), 1.88-2.02(br. π , 4H), 2.3(d, 1H), 2.35(m, 1H), 2.45(m, 1H), 2.55-2.75(π , 3H), 3.0(s, 6H), 3.25(π , 1H), 3.55(π , 1H), 3.65(π , 1H),
- 20 3.75(m, 1H), 3.9(m, 1H), 4.3(t, 1H), 4.55(m, 2H), 4.68(br. m, 1H), 3.9(m, 1H), 4.3(t, 1H), 4.55(m, 2H), 4.68(br. m, 1H), 4.95(br. m, 1H), 5.1(br. m, 2H), 5.45(d, 1H), 6.5(m, 2H), 7.7(m, 2H).

[1S,9S(2RS,3S)]9-[(3-Chloro-4-aminobenzoyl)amino]-6,10-

- 25 dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-ethoxy-5oxotetrahydrofuran-3-yl)-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (550h),
 was synthesized from 212h by the methods used to
 prepare 213e from 212e to afford 195 mg of 550h as a
- 30 white solid, ^{1}H NMR (DMSO-d₆) δ 1.1-1.18(2t, 3H), 1.6-

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[1S,9S(2RS,3S)]9-[(4-Methoxybenzoyl)amino]-6,10-dioxo-

- 10 1,2,3,4,7,8,9,10-octahydro-N-(2-ethoxy-5-oxotetrahydrofuran-3-v1)-6H
 - pyridazino[1,2-a][1,2]diazepine-1-carboxamide (5501),
 was synthesized from 212i by the methods used to
 prepare 213e from 212e to afford 135 mg of 550i, ¹H NMR
- 15 (CDC1₃) δ 1.18-1.28(2t, 3H), 1.6-1.75(m, 1.5H), 1.9-2.1(m, 3.5H), 2.22-2.3(d, 0.5H), 2.38-2.47(m, 1.5H), 2.7-2.8(m, 0.5H), 2.8-2.93(m, 1H), 2.94-3.15(m, 1.5H), 3.15-3.28(m, 1H), 3.55-3.62(q, 0.5H), 3.62-3.73(q, 0.5H), 3.78-3.88(α, 0.5H), 3.88(α, 3H), 3.9-3.95(α,
- 20 0.5H), 4.33-4.4(m, 0.5H), 4.5-4.55(m, 1H), 4.68-4.76(m, 0.5H), 4.9-4.95(m, 0.5H), 5.1-5.2(m, 1.5H), 5.18(s, 0.5H), 5.48-5.52(d, 0.5H), 6.48-6.55(d, 0.5H), 6.85-6.9(m, 1H), 6.9-6.95(m, 2H), 7.34-7.38(d, 0.5H), 7.78-7.85(m, 2H).
- 25 [18,98(2RS,38)]9=[(3,5-Dichloro-4hydroxybenzoyl)amino]-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-N-(2-ethoxy-5-oxotetrahydrofuran-3-yl)-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (550k), was synthesized from 212k by the methods used to prepare 213e from 212e to afford 174 ng of 550k as a
- 30 prepare **213e** from **212e** to afford 174 mg of **550k** as white solid, 1 H NMR (DMSO-d₆) δ 1.15(2t, 3H), 1.6-

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1.75 (m, 2H), 1.9-2.05 (m, 2H), 2.1-2.4 (m, 5H), 2.5-2.55 (m, 1H), 2.7-2.8 (m, 0.5H), 2.85-3.0 (m, 1H), 3.0-3.1 (m, 0.5H), 3.55-3.7 (m, 1H), 3.7-3.8 (m, 1H), 4.2 (t, 0.5H), 4.35-4.45 (m, 0.5H), 4.55-4.65 (m, 0.5H), 4.8-5 (m, 0.5H), 5.05 (t, 0.5H), 5.15 (t, 0.5H), 5.35 (s, 0.5H), 5.6 (d, 0.5H), 7.95 (s, 2H), 8.5 (d, 0.5H), 8.65 (d, 1H), 8.75 (d, 0.5H), 10.9 (br. s, 1H).

[15,95(2RS,3S)]9-[(3-Chloro-4-acetamidobenzoy1)amino]-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-ethoxy-5-

- 15 2.15(m, 3.5H), 2.22-2.28(m, 0.5H), 2.28(s, 3H), 2.38-2.48(m, 1.5H), 2.66-2.92(m, 1.5H), 2.95-3.14(m, 1.5H), 3.2-3.34(m, 1H), 3.56-3.63(q, 0.5H), 3.63-3.72(q, 0.5H), 3.8-3.85(q, 0.5H), 3.9-3.95(q, 0.5H), 4.32-4.38(m, 0.5H), 4.5-4.62(m, 1H), 4.68-4.75(m, 0.5H),
- 20 4.88-4.92(m, 0.5H), 5.08-5.2(m, 1.5H), 5.18(s, 0.5H), 5.46-5.5(d, 0.5H), 6.5-6.55(d, 0.5H), 6.98-7.05(m, 1H), 7.42-7.48(d, 0.5H), 7.63-7.78(m, 2.5H), 7.9-7.94(d, 0.5H), 8.44-8.52(m, 1H).

[1S,9S(2RS,3S)]9-[(3,5-Dichloro-4-

25 methoxybenzoyl)amino]-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-N-(2-ethoxy-5-oxotetrahydrofuran-3-yl)-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (550m),
was synthesized from 212m by the methods used to
prepare 213e from 212e to afford 301 mg of 550m as a
30 white solid, ¹H NMR (CDCl₃) & 1.2-1.35(2t, 3H), 1.5-

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1.8 (m, 2H), 1.9-2.15 (5H), 2.25 (d, 0.5H), 2.4-2.5 (m, 2H), 2.65-2.8 (m, 0.5H), 2.8-3.0 (m, 0.5H), 3.0-3.2 (m, 1F), 3.2-3.35 (m, 0.5H), 3.55-3.65 (m, 0.5H), 3.65-3.75 (m, 0.5H), 3.8-3.9 (m, 0.5H), 3.9-4.0 (m, 0.5H), 4.4-4.45 (m, 0.5H), 4.55-4.65 (m, 0.5H), 4.7-4.8 (m, 0.5H), 4.85-4.95 (m, 0.5H), 5.05-5.2 (m, 0.5H), 5.2 (s, 0.5H), 5.5 (d, 0.5H), 6.5 (d, 0.5H), 6.9 (d, 0.5H), 6.95 (d, 0.5H), 7.35 (d, 0.5H), 7.75 (s, 1H), 7.85 (s, 1H).

[3S(1S,9S)]3-(9-(3,5-Dichlorobenzoyl)amino-6,10-dioxo-10 1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (214j), was synthesized from 213j by the method used to prepare 2002 from 2001 to afford 62 mg of 214j as a white solid, $^{1}{\rm H~NMR~(CD_3OD)}~\delta$ 0.9 (t.

- 15 1H), 1.3(br. s, 1H), 1.7(br. m, 1H), 1.9(br. m, 1H), 2.1(br. s, 1H), 2.25(q, 1H), 2.35(m, 1H), 2.48(m, 2H), 2.65(t, 1H), 3.15(br. t, 1H), 3.5(br. m, 1H), 4.3(br. s, 1H), 4.55(m, 2H), 4.95(t, 1H), 5.25(br. s, 1H), 7.6(br. s, 1H), 7.85(br. s, 1H)
- 20 [3S(1S,9S)]3-(9-(3,5-Dichloro-4-hydroxybenzoy1) amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (214k), was synthesized from 213k by the method used to prepare 2002 from 2001 to afford £0 mg of 214k as a white solid, ¹H NMR (CD₃OD) δ 1.6-1. ⁷ m, 1H), 1.8-2.0 (m, 2H), 2.0-2.1 (m, 2H), 2.15-2.25 (m, 1H), 2.3-2.4 (m, 1H), 2.4-2.55 (m, 2H), 2.6-2.75 (m,1H), 3.05-3.2 (m, 1H), 3.4-3.6 (m, 2H), 4.2-4.3 (m, 1H), 4.45-4.6 (m, 1H), 4.8-5.0 (m, 1H), 5.1-5.2 (m, 1H), 7.85 (s, 2H).

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[3S(1S,9S)]3-(9-(3-Chloro-4-acetamidobenzoy1)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (2141), was synthesized from 2131 by the method used to prepare 2002 from 2001 to afford 91 mg of 2141 as a white solid, ¹H NMR (DMSO-d₆) & 1.65(br.m, 6H), 1.9(br.m, 6H), 2.15(s, 3H), 2.3(m, 3H), 2.6-2.85(m, 3H), 2.9(m, 2H), 3.0(m, 1H), 4.15(br.q, 1H), 4.4(m, 3H), 5.0(m, 1H), 5.15(m, 1H), 5.45(s, 1H), 7.8(d, 2H), 7.95(d, 1H), 8.05(s, 1H), 8.65(m, 2H),

[3s(1s,9s)]3-(9-(3,5-Dichlorobenzoyl)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-

9.65(s. 1H).

- oxobutanoic acid (214m), was synthesized from 213m by the method used to prepare 2002 from 2001 to afford 105 mg of 214m as a white solid, $^1\mathrm{H}$ NMR (CD_3OD) δ 1.6-1.75(m, 1H), 1.85-1.95(m, 1H), 2.0-2.1(m, 2H), 2.15-2.25(m, 1H), 2.3-2.4(m, 1H), 2.45-2.55(m, 2H), 2.65-
- 20 2.75(m, 1H), 3.4-3.55(m, 2H), 3.95(s, 3H), 4.2-4.3(m, 1H), 4.45-4.6(m, 1H), 4.9-5.0(m, 1H), 5.15-5.2(m, 1H), 7.9(s, 2H).

Compounds $\mathbf{308c}$ and $\mathbf{308d}$ were prepared as follows.

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[3S(1S,9S) 3-(9-(4-Methoxybenzoyl)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-amino]-4-oxobutanoic acid, O-methyl oxime (308c), was

- 5 synthesized from 212e via the methods used to prepare 308b from 212e to afford 266 mg of 308c 1 H NMR (CDCl $_{3}$) δ 1.6-1.7 (m, 1H), 1.88-1.98 (m, 3H), 2.02-2.15 (m, 1H), 2.3-2.4 (m, 1H), 2.65-2.95 (m, 3H), 3.04-3.09 (m, 1H), 3.12-3.25 (m, 1H), 3.84 (s, 3H), 3.86 (s, 3H), 4.5-4.58 (m, 1H), 4.88-4.95 (m, 1H), 5.1-5.25 (m, 2H), 6.86-6.9 (d,
- 10 1H), 4.88-4.95(m, 1H), 5.1-5.25(m, 2H), 6.86-6.9(d, 2H), 7.15-7.25(m, 2H), 7.36-7.4(m, 1H), 7.75-7.8(d, 2H).

[3S(1S,9S) 3-(9-(4-Methoxybenzoy1) amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

- - 4.88-4.95(m, 1H), 5.05(s, 2H), 5.1-5.2(m, 1H), 6.82-

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6.95(m, 2E), 7.02-7.15(m, 2H), 7.28(m, 5H), 7.45(m, 1H), 7.72(d, 2H).

Compounds 2100f, 2100g, 2100h, 2100i and 2100j were prepared as described below.

5 (3S,2RS) 3-Allyloxycarbonylamino-2-(4-chlorobenzyl)oxy-5-oxotetrahydrofuran (2101a), was synthesized from allyloxycarbonylamino-β-tert-butyl aspartate by the methods employed by Chapman (Ricorg. & Med. Chem. Lett., 2, pp.615-618 (1992)) to prepare (3S,2RS) 3allyloxycarbonylamino-2-benzyloxy-5-oxotetrahydrofuran

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using 4-chlorobenzyl alcohol instead of benzyl alcohol to afford 1.84 q of 2101a as a crystalline solid.

[15.95(2RS.3S)] 9-Benzovlamino-6.10-dioxo-

1,2,3,4,7,8,9,10-octahydro-N-(2-(4-chlorobenzyl)oxy-55 oxotetrahydrofuran-3-vl)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (2100f), was synthesized from 212e by the methods used to prepare 213e from 212e using 2101a to afford 380 mg of 2100f, 1 H NMR (CDCl₃) δ 1.8-2.0 (m, 10H), 2.30 (d, 1H),

10 2.31-2.5(m, 3H), 2.7-2.9(m, 3H), 3.05(m, 2H), 3.1-3.2(m, 4H), 4.45(q, 1H), 4.5-4.6(m, 3H), 4.7(d, 2H), 4.85(d, 1H), 4.9(t, 1H), 5.2(t, 1H), 5.15(m, 2H), 5.25(s, 1H), 5.55(d, 1H), 6.5(d, 1H), 6.9(d, 1H), 6.95(d, 1H), 7.25(m, 3H), 7.35(t, 2H), 7.45(m, 2H),

15 7.55(1H), 7.8(m, 3H).

(3S,2RS) 3-Allyloxycarbonylamino-2-anti-isopropoxy-5oxotetrahydrofuran (2101b), was synthesized from
(3S,2RS) 3-allyloxycarbonylamino-2-benzyloxy-5oxotetrahydrofuran via the method used to prepare 2100d
20 from 214e using H₂SO₄ instead of pTSA to afford 2101b.

[1s,9s(2Rs,3s)] 9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-anti-isopropoxy-5oxotetrahydrofuran-3-yl)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (2100g),

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4.52-4.63 (m), 4.90-4.95 (m), 5.12-5.20 (m), 5.28 (s), 6.93 (d), 7.10 (d), 7.41-7.50 (m), 7.51-7.58 (m), 7,84 (d).

[1s,9s(2Rs,3Rs)] 9-Benzoylamıno-6,10-dioxo-

5 1,2,3,4,7,8,9,10-octahydro-N-(2-acetoxy-5-oxotetrahydrofuran-3-y1)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (2100h).
A solution of 214e (287 mg, 0.65 mmol) in pyridine (5
mL) was treated with Ac₂O (0.4 mL, 3.62 mmol). After 6

- 10 hours, the reaction mixture was poured into 5% NaHSO4 and extracted 3 times with EtOAc. The combined organics were washed with brine, dried over Na₂SO₄ and concentrated in vacuo. Chromatography (SiO₂, EtOAc) afforded 119 mg of 2100h, ¹HNMR (CDCl₂, mixture of four
- 15 diastereoisomers) δ 1.80-2.05(m), 2.12(s), 2.13(s), 2.19(s), 2.22(d), 2.67-2.75(m), 2.80-2.95(m), 3.00-3.20(m), 3.21-3.33(m), 3.50-3.95(four discrete multiplets), 4.19(m), 4.55(m), 4.57-4.65(m), 4.69(m), 4.85-4.95(m), 5.04(m), 5.10(s), 5.10-5.22(m), 6.46(d),
- 20 6.03(s), 6.50(d), 6.58(d), 6.75(d), 6.95-7.05(m), 7.22(m), 7.30(m), 7.71(d), 7.75-7.83(m).

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[35(15,95)]3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid ethyl ester (2100i). To a solution of 2100b (1.5 g, 2.7 mmol) in CH₃CN (10 mL) was added 1N HCl at ambient temperature. After 6 hours solid NaHCO₃ was added and the product extracted with EtoAc, dried over MgSO₄ and concentrated in vacuo. Chromatography (SiO₂, 30-100% CH₂Cl₂ in EtoAc) afforded 123 mg of

10 2100i, ¹H NMR (CDCl₃) δ 1.25(t, 3H), 1.6-1.8(m, 1H), 1.9-2.2(m, 5H), 2.4-2.5(m, 1H), 2.75-2.9(m, 2H), 3.0-3.1(m, 2H), 3.2-3.25(m, 1H), 4.05-4.2(m, 1H), 4.5-4.7(m, 1H), 5.1-5.25(m, 1H), 7.0-7.2(m, 2H), 7.4-7.45(m, 2H), 7.5(t, 1H), 7.8(t, 2H), 9.5(s, 1H).

15 [3s(1s,9s)]3-(9-Benzoylamino-6,10-dioxo1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4acetoxy-3-butenoic acid ethyl ester (2100j), was
synthesized from 2100i via the method used to prepare

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2100h from 214e to afford 347 mg of 2100, ¹H NMR (CDCl₃) & 1.3(t, 3H), 1.6-1.8(m, 2H), 1.9-2.25(m, 4H), 2.25(s, 3H), 2.3-2.45(m, 1H), 2.8-3.0(m, 1H), 3.0-3.25(m, 2H), 3.4-3.45(m, 2H), 4.1-4.2(m, 2H), 4.55-5 4.7(m, 1H), 5.1-5.25(m, 1H), 6.8(s, 1H), 7.0-7.1(m, 2H), 7.5(t, 1H), 7.8(t, 2H), 9.5(s, 1H).

Compounds 500 and 501 are described in Table 23. These compounds were prepared by methods similar to the methods used to prepare compounds 404-449 (see, 10 Example 11).

+ (H+W) 533 MS HPLC RT min 10.13 0.97 11.448 (A) (method) Purity 0.991 C22H24C1N508 | 521.92 532.51 Σ C24H28N4O10 MF Structure Compound 500 501

Table 23

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The compounds described below (213m, 213n, 213o, 213p, 213q, 213r, 213s, 213t, 213u, 213v, 213w, 213x, and 214w), were prepared by methods similar to the methods used to prepare compounds 213b-f.

Compounds **419**, **415**, **450**, **456**, **475**, **404**, **486**, **487**, **417**, **408** and **418** may also be prepared as described below.

213m-x 214w, 404, 408, 415,

417, 418, 419, 450,

15 456, 475, 486, 487

10

compound	R ¹	
213m, 419	MeOC(O)-	
213n, 415		

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2130, 450	o in the second
213p, 456	но
213q, 475	S) NH
213r, 404	ו • • • • • • • • • • • • • • • • • •
213s, 486	
213t, 4 87	
213u, 417	MeO HeO

5

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213v, 408	
213w, 214w	Me HO Me
213x, 418	H ₃ C H _N

[15,98(2R5,35)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-5 yl)-6,10-dioxo-9-(3,4-methylenedioxybenzoylamino)-

1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213n),

was isolated as a mixture of diastereomers (syn:anti isomer ratio 6:4) (1.43g, 82%) as a white solid: mp.

- 10 206-10°C; IR (KBr) 3288, 1787, 1680, 1657, 1651, 1619, 1548, 1440, 1256, 1135;

 1 M NMR (D₆-DMSO) δ 8.75 (0.4H, d), 8.55 (0.6H, d), 8.45 and 8.43 (1H, 2 x d), 7.50 (1H, d), 7.42 (1H, s), 7.40-7.27 (5H, m), 7.01 (1H, d), 6.11 (2H, s), 5.67 (0.6H, d), 5.43 (0.4H, s), 5.10-5.09
- 15 (1H, m), 4.90-4.59 (3.5H, m), 4.45-4.25 (1.5H, m), 3.47-3.20 (1H, m), 3.20-2.70 (2H, m), 2.65-2.35 (1H, m), 2.35-2.00 (3H, m), 2.00-1.75 (2H, m), 1.65-1.40 (2H, m). Anal. Calcd for C₂₉H₃₀N₄O₉: C, 60.20; H, 5.23; N, 9.68. Found: C, 60.08; H, 5.32; N, 9.50. MS (ES⁷)

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580 $(M^+ + 2, 35\%)$, 579 $(M^+ + 1, 100)$, 404 (5), 367 (5), 236 (7), 107 (5).

[15,95(2R5,35)]9-[(3-Acetamido)benzamido]-N-(2-benzyloxy-5-oxotetrahydrofuran-3-y1)-6,10-dioxo-

5 1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (2130), anti-isomer as a white foamy solid (0.73g, 69%): mp. $135-40^{\circ}\mathrm{C}$; [α] $_{\mathrm{D}}^{21}$ -37.3° (c 0.1, CH₂Cl₂); IR (KBr) 3452, 3310, 1790, 1664, 1659, 1650, 1549, 1425, 1258, 1121;

- 10 1 H NMR (D₆-DMSO) δ 10.11 (1H, s), 8.77 (1H, d), 8.57 (1H, d), 8.01 (1H, s), 7.76 (1H, d), 7.55 (1H, d), 7.45-7.25 (6H, m), 5.43 (1H, s), 5.08-5.00 (1H, m), 4.95-4.73 (1H, m), 4.76 and 4.68 (2H, dd), 3.40-3.20 (1H, m), 3.09 (1H, dd), 3.02-2.75 (1H, m), 2.45-2.06
- 15 (4H, m), 2.06 (3H, s), 2.00-1.75 (2H, m), 1.70-1.40 (2H, m). Anal. Calcd for $C_{30}H_{33}N_{5}O_{8}^{+}0.75H_{2}O$: C, 59.54; H, 5.75; N, 11.57. Found: C, 59.40; H, 5.62; N, 11.50. MS (ES⁴) 593 (M⁴ + 2, 33%), 592 (M⁴ + 1, 100), 574 (7), 487 (7), 475 (6), 385 (9), 373 (26), 318 (14), 296 (11), 266 (10), 221 (22).

 $[1s,9s(2Rs,3s)] \ \ N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(4-hydroxybenzoyl)amino-$

1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213p),

25 was isolated as a foam (1.2g, 77%): $[\alpha]_0^{20}$ -115° (c 0.20, CH₂Cl₂); IR (KBr) 3368, 2946, 1794, 1654, 1609, 1540, 1505, 1421, 1277, 1175, 1119, 980; 1 H NMR (D₆-DMSO) δ 10.1 (1H, s), 8.80 (0.5H, d, J = 6.6), 8.60 (0.5H, d, J = 7.2), 8.40-8.36 (1H, 2d), 7.82 (2H, d, J 30 = 8.0), 7.41 (5H, bs), 6.86 (2H, d, J 8.6), 5.72 (0.5H,

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d, J = 5.0), 5.49 (0.5H, bs), 5.13-5.07 (1H, m), 4.95-4.65 (2.5H, m), 4.49-4.38 (2.5H, m), 3.49-3.30 (2H, m), 3.21, 2.79 (2H, m), 2.40-1.41 (7H, m). MS (ES⁺) 551.

[1S,9S(2RS,3S)]N-(2-Benzyloxy-5-oxotetrahydrofuran-3-5 yl)-6,10-dioxo-9-(indol-2-oylamino)-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2)diazepine-1carboxamide (213q), was isolated as a white glassy solid (80%): mp. 145-149°C; [α]_D²³ -56.0° (c 0.05, CH₂Cl₂); IR (KBr) 3399-3319, 1791, 1657, 1543, 1420, 1253, 1119; ¹H NMR (CDCl₃) 89.54 (1H, s), 7.65 (1H, d, J = 7.9), 7.51 (1H, d, J = 6.9), 7.44-7.25 (7H, m), 7.18-7.06 (3H, m), 5.30-5.20 (1H, m), 5.27 (1H, s), 4.84 (1H, m), 4.79 (1H, d, J = 11.4), 4.56 (1H, d, J = 11.3), 4.47 (2H, m), 3.28 (1H, m), 3.10-2.97 (2H, m),

15 2.71 (1H, m), 2.47-2.37 (1H, m), 2.26 (1H, d, J = 17.9), 2.09 (1H, m), 1.83, 1.70, 1.51 (4H, 3m).

[1s,9s(2Rs,3s)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-9-(2-tolucylamino)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213r), was isolated as a mixture of diastereomers (syn:anti isomer ratio 55:45) as a white foamy solid (1.46g, 89%): mp. 106-10°C; IR (KBr) 3306, 2947, 1791, 1659, 1650, 1535, 1421, 1256, 1122; ¹H NMR (D₆-DMSO) & 8.76 (0.45H, d), 8.56 (0.55H, d), 8.49 and 25 8.47 (1H, 2 x d), 7.41-7.19 (9H, m), 5.67 (0.55H, d), 5.43 (0.45H, s), 5.11-5.02 (1H, m), 4.86-4.55 (3.5H, m), 4.45-4.25 (1.5H, m), 3.40-3.20 (1H, m), 3.20-2.70 (2H, m), 2.65-2.40 (1H, m), 2.34 (3H, s), 2.30-1.70 (5H, m), 1.65-1.40 (2H, m). Anal. Calcd for C29H37NgO9:

30 C, 62.66; H, 5.95; N, 10.08. Found: C, 62.91; H, 6.00;

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N, 9.70. MS (ES^+) 550 $(M^+ + 2, 43\%)$, 549 $(M^+ + 1, 100)$, 374 (3), 280 (4), 279 (20), 118 (5).

[1S,9S(2RS,3S)]N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-9-[4-

5 (phenylacetamido) benzamido] -6H-

pyridazino[1,2-a][1,2]diazepin-1-carboxamide (213s), was isolated as the anti-isomer as a white foamy solid (0.64g, 77%): mp. 137-41°C; $\left[\alpha\right]_D^{21}$ -48.2° (c 0.05, CH₃OH); IR (KBr) 3477, 3314, 1791, 1659, 1599, 1529,

10 1499, 1406, 1256, 1122; ¹H NMR (D₆-DMSO) & 10.45 (1H, s), 8.76 (1H, d), 8.50 (1H, d), 7.86 (2H, d), 7.69 (2H, d), 7.41-7.20 (10H, m), 5.43 (1H, s), 5.08-4.98 (1H, m), 4.90-4.73 (1H, m), 4.76 and 4.68 (2H, dd), 3.67 (2H, s), 3.40-3.20 (1H, m), 3.09 (1H, dd), 3.02-2.75

15 (1H, m), 2.39 (1H, dd), 2.30-2.00 (3H, m), 2.00-1.75 (2H, m), 1.70-1.40 (2H, m). Anal. Calcd for C₃₆H₃₇N₅C₉·0.5H₂O: C, 63.90; H, 5.66; N, 10.35. Found: C, 63.68; H, 5.67; N, 10.24. MS (ES[†]) 669 (M[†] + 2, 40%), 668 (M^{*} + 1, 100), 640 (12), 435 (18), 425 (23), 20 403 (33), 328 (17), 302, (32), 274 (22), 197 (16), 138 (17).

 $\label{eq:continuous} $$ [1S,9S(2RS,3S)]$ N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-[4-(3-methylbutan-1-oylamino)benzamido]-1,2,3,4,7,8,9,10-octahydro-6H-$

25 **pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213t)**, was isolated as a white foamy solid (0.63g, 80%,: mp. $159-64^{\circ}C$; [α]₀ 21 -37.0° (c 0.05, CH₃OH); TR (KBr; 3463, 3321, 1790, 1680, 1658, 1650, 1644, 1595, 1525, 1501, 1408, 1251, 1113, 933; 1 H NMR (D_{6} -DMSO) δ 10.13 | H, s;, 30 8.76 (1H, d), 8.48 (1H, d), 7.85 (2H, d), 7.68 (2H, d),

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 $7.40-7.25 \ (5H, m), \ 5.43 \ (1H, s), \ 5.08-4.95 \ (1H, m), \\ 4.92-4.73 \ (1H, m), \ 4.76 \ and \ 4.68 \ (2H, dd), \ 3.40-3.20 \\ (1H, m), \ 3.09 \ (1H, dd), \ 3.02-2.75 \ (1H, m), \ 2.39 \ (1H, dd), \ 2.35-2.00 \ (6H, m), \ 2.00-1.75 \ (2H, m), \ 1.70-1.40$

- 10 [1S,9S(2RS,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3yl)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-9-(3,4,5trimethoxybenzoylamino)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213u),
 was isolated as a white solid (81%): mp. 120-132°C; IR
15 (KBr) 3361-3334, 1792, 1659, 1585, 1536, 1499, 1457,

- 1416, 1340, 1236, 1126, 989; 1 H NMR (CDC1₃) δ 7.39-7.29 (6H, m), 7.12 (1H, s), 7.03 (1H, s), 6.92, 6.83, 6.48 (approx 3H, 3d, J = 8.1, 7.5, 8.1), 5.57 (d, J = 5.3), 5.27 (1H, s), 5.23-5.06, 4.91-4.71, 4.64-4.43, (6H,
- 20 3m), 3.92, 3.91, 3.89, 3.88 (9H, 4s), 3.32-2.70, 2.52-2.08, 1.91, 1.63 (1H, 4m).

[1s,9s(2rs,3s)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-y1)-6,10-dioxo-9-(naphth-1-oylamino)-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-

25 **carboxamide (213v)**, was isolated as a white solid (78%): mp. 121-7°C; IR (KBr) 3534-3331, 1791, 1659, 1528, 1420, 1256, 1122; ¹H NMR (CDCl₃) & 8.34-8.29 (1H, m), 7.98-7.87 (2H, m), 7.68-7.45 (4H, m), 7.34-7.24 (5H, m), 7.04 (d, J = 6.8), 6.78 (d, J = 7.8), 6.66 (d, 30 J = 7.7), 6.48 (2H, d, J = 7.5)5.56 (d, J = 5.4), 5.10

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(1H, s), 5.30-5.14, 5.0, 4.89 (d, J = 11.2), 4.71-4.41 (6H), 3.18-2.80, 2.50-2.27, 2.08-1.60 (11H, 3m).

[15,95(2R5,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(4-hydroxy-3,5-dimethylbenzoyl)amino-5 1,2,3,4,7,8,9,10-octahydro-6H-

- pyridazino[1,2-a][1,2]diazepine-1-carboxamide (213w),
 was isolated as a mixture of diastereoisomers (65/35)
 as a white solid (0.9g, 65%): mp. 110-115°C (decomp.);
 IR (KBr) 3409, 2945, 1792, 1658, 1606, 1534, 1486,
- 10 1420, 1330, 1276, 1209, 1122, 980, 960; ¹H NMR (CDC1₃) δ 7.66 (0.35H, d, J = 6.9), 7.46-7.20 (7H, m), 6.93 (0.35H, d, J = 7.7), 6.85 (0.65H, d, J = 7.6), 6.73 (0.65H, d, J = 7.6), 5.96 (0.35H, bs), 5.85 (0.65H, bs), 5.56 (0.65H, d, J = 5.2), 5.28 (0.35H, bs), 5.56 (0.65H, d, J = 5.2), 5.28 (0.35H, bs), 5.20-
- 15 4.98 (2H, m), 4.96-4.40 (4H, m), 3.28-2.55 (3H, m), 2.53-2.32 (1H, m), 2.23 (6H, 2s), 2.03-1.40 (7H, m). MS (ES⁻) 577, (ES⁺) 579.

[15,95(2R5,35)] 9-[4-(Acetylamino)benzoylamino]-N-(2-benzyloxy-5-oxo-tetrahydrofuran-3-yl)-6,10-dloxo-

- 20 1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboximide (213x),
 was isolated as a colourless poweder (691mg, 86%): mp.
 150-70°C; [α]_D²² -10.1° (c 0.10, Me₂CO); IR (KBr) 3313,
 1791, 1679, 1654, 1597, 1528, 1501, 1457, 1407, 1371,
- 25 1315, 1255, 1184, 1122, 933; ¹H NMR (d6-DMSC) & 8.75 (1H, d1, 8.47 (1H, d), 7.84 (2H, d), 7.66 (2H, d), 7.35 (5H, m1, 5.43 (1H, s), 5.06-5.03 (1H, m), 4.90-4.64 (3H, m), 4.46-4.26 (2H, m), 3.16-2.86 (2H, m), 2.45-2.05 (5H, m1, 2.07 (3H, s), 2.00-1.84 (2H, m), 1.68-1.56 (2H, m);
- 30 Anal. Calcd for $C_{30}H_{33}N_5O_8 \cdot H_2O$: C, 59.11; H, 5.79; N,

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11.49. Found: C, 59.38; H, 5.66; N, 11.31; M.S. (ES^{+}) 614 (100%), 592 $(M^{+}+1.66)$.

[3S(1S,9S)] 3-[6,10-Dioxo-9-(3,4-

methylenedioxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-

- 5 6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxobutanoic acid (415), was prepared by a similar method as compound 214e to afford a white solid (297mg, 84%): mp. 158-62°C; [α]_D²⁴ -109.5° (c 0.1, CH₃OH); IR (KBr) 3700-2500 (br), 1783,1659, 1650, 1538, 1486,
- 10 1439, 1257, 1037; ^{1}H NMR (CD₃OD) δ 7.48 (1H, dd), 7.35 (1H, d), 6.88 (1H, d), 6.03 (2H, s), 5.25-5.15 (1H, m), 5.02-4.90 (1H, m), 4.63-4.45 (2H, m), 4.30-4.20 (1H, m), 3.57-3.30 (1H, m), 3.20-3.05 (1H, m), 2.75-2.10 (5H, m), 2.10-1.60 (4H, m). MS (ES †) 488 (M+, 25%),
- 15 487 (M^+ 1, 100), 443 (8), 387 (3), 315 (5), 150 (6), 127 (5), 113 (8). Accurate mass calculated for $C_{22}H_{25}N_4O_9$ (MH^+): 489.1621. Found 489.1646.

[3S(1S,9S)] 3-{9-[(3-Acetamido)benzamido]-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

- 20 pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4 oxobutanoic acid (450), was prepared by a similar
 method as compound 214e to afford a white foamy solid
 (378mg, 94%): mp. 175-9°C; (α)_D²² -91.7° (c 0.1, CH₃OH);
 IR (KBr) 3700-2500 (br), 3319, 1659, 1590, 1553, 142″,
- 25 1260; ¹H NMR (CD₃OD) & 8.01 (1H, d), 7.74 (1H, dd), 7.56 (1H, d), 7.45-7.35 (1H, m), 5.25-5.15 (1H, m), 5.05-4.90 (1H, m), 4.60-4.45 (2H, m), 4.30-4.20 (1H, m), 3.55-3.30 (1H, m), 3.20-3.00 (1H, m), 2.75-2.20 (5H, m), 2.14 (3H, s), 2.20-1.60 (4H). Anal. Caicd for
- 30 $C_{23}H_{27}N_5O_8 \cdot 1.5H_2O$: C, 52.27; H, 5.72; N, 13.25. Found:

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C, 52.31; H, 5.86; N, 12.85. MS (ES^+) 501 $(M^+, 26\%)$, 500 $(M^+ - 1, 100)$, 328 (2), 149 (3), 113 (3).

[3S(1S,9S)] 3-[4-(Hydroxybenzoyl)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

- 5 pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4oxobutanoic acid (456), was prepared by a similar
 method as compound 214e to afford a white solid (0.73g,
 72%): mp. >260°C; [α]_D²⁰ -66° (c 0.34, MeOH); IR (KBr)
 3401, 2946, 1651, 1609, 1584, 1506, 1426, 1277, 1257,
- 10 1177; 1 H NMR (D_{6} -DMSO) δ 10.2 (1H, very bs), 9.17 (1H, bs), 8.65 (1H, s), 8.37 (1H, d, J 5.4), 7.81 (2H, d, J = 8.2), 6.87 (2H, d, J = 8.4), 5.24 (1H, m), 4.92-4.86 (1H, m), 4.41-4.32 (2H, m), 3.68-3.21 (3H, m), 3.12-2.79 (1H, m), 2.50-1.42 (7H, m). MS (ES $^{+}$) 459.
- 15 [3S(1S,9S)] 3-[6,10-Dioxo-9-(indol-2-oylamino)1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4oxobutanoic acid (475), was prepared by a similar
 method to that described for compound 214e to afford a
 20 white solid (79%): mp. 150°C (softens) 190-210°C;
 - white solid (79%): mp. 150°C (softens) 190-210°C; $[a]_D^{23} -97.5^{\circ} (c 0.1, CH_3OH); IR (KBr) 3319, 1658, 1650, 1549, 1421, 1256; ^1H NMR (CD_3OD) & 7.61 (1H, d, J = 8.0), 7.43 (1H, d, J = 8.1), 7.21 (2H, m), 7.05 (1H, m), 5.21 (1H, m), 5.07-4.77 (1H, m), 4.54 (2H, m), 4.23 (1H, m), <math> ...$
- 25 3.46 (1H, m), 3.14 (1H, m), 2.66-1.71 (9H, m). MS (ES^{*}, m/z), 482 (M^{*} 1, 100%).

[35(15,95)] 3-[6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-9-(2-toluoylamino)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxobutanoic acid (404), was prepared by

a similar method as compound **214e** to afford a white solid (0.79g, 86%): mp. 156-9°C; $[\alpha]_D^{25}$ -119.7° (c 0.1, CH₃OH); IR (KBr) 3700-2500 (br), 3387, 3309, 2956, 1785, 1659, 1650, 1535, 1422, 1278; ¹H NMR (CD₃CD) δ 5 7.46-7.15 (4H, m), 5.25-5.15 (1H, m), 5.02-4.90 (1H, m), 4.58-4.45 (2H, m), 4.30-4.20 (1H, m), 3.55-3.30 (1H, m), 3.20-3.05 (1H, m), 2.80-2.20 (4H, m), 2.41 (3H, s), 2.20-1.60 (5H, m). MS (Ξ S^{*}) 458 (M+, 27%), 457 (M[†] - 1, 100), 413 (13), 339 (8), 285 (5), 134 (6), 10 127 (11). Accurate mass calculated for C₂₂H₂₇N₄O₇ (MH[†]): 459.1880. Found 459.1864.

[3S(1S,9S)] 3-(6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-9-[4-(phenylacetamido)benzamido]-6Hpyridazino[1,2-a][1,2]

- 15 diazepine-1-carboxamido}-4-oxobutanoic acid (486), was prepared by a similar method as compound 214e to afford a white solid (325mg, 89%): mp. 165-9°C; [α]_D²² -69.1° (c 0.1, CH₃OH); IR (KBr) 3700-2500 (br), 3318, 1658, 1599, 1530, 1505, 1407, 1258; ¹H NMR (CD₃OD) δ 7.85 (2H, 2d), 7.69 (2H, d), 7.38-7.20 (5H, m), 5.25-5.15 (1H, m), 5.05-4.90 (1H, m), 4.57-4.45 (2H, m), 4.30-4.20 (1H, m), 3.70 (2H, s), 3.55-3.30 (1H, m), 3.20-3.00 (1H, m), 2.75-1.60 (9H, m). Anal. Calcd for C₂dH₃N₅O₈·1.5H₅O:
- C, 57.61; H, 5.67; N, 11.58. Found: C, 57.81; H, 5.74; 25 N, 11.47. MS (ES^{+}) 577 $(M^{+}, 33^{\circ})$, 576 $(M^{+} 1, 100)$, 502 (2).

[3S(1S,9S)] 3-{6,10-Dioxo-9-{4-(3-methylbutan-1-oylamino)benzamido]-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-30 oxobutanoic acid (487), was prepared by a similar

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method as compound 214e to afford a white foamy solid (335mg, 93%): mp. 176-80°C; $[\alpha]_D^{22}$ -88.0° (c0.1, CH₃OH); IR (KBr) 3700-2500 (br), 3321, 2960, 1781, 1660, 1597, 1529, 1407, 1258, 1167; ¹H NMR (CD₃OD) δ 7.86 (2H, d), 7.69 (2H, d), 5.25-5.15 (1H, m), 5.05-4.90 (1H, m), 4.60-4.45 (2H, m), 4.30-4.20 (1H, m), 3.57-3.30 (1H, m), 3.20-3.00 (1H, m), 2.75-1.60 (12H, m), 1.00 (6H, d). Anal. Calcd for C₂GH₃3N₅O₈·H₂O: C, 55.61; H, 6.28; N, 12.45. Found: C, 56.00; H, 6.37; N, 12.15. MS 10 (ES[†]) 543 (M+, 31%), 542 (M[†] - 1, 100), 498 (2), 468 (3).

[3S(1S,9S)] 3-[6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-9-(3,4,5-trimethoxybenzoylamino)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-

- 15 oxobutanoic acid (417), was prepared by a similar
 method to that described for compound 214e to afford a
 white solid (0.63g, 92%): mp. 145-155°C (approx., not
 sharp); [α]_D²⁷ -114.6° (c 0.11, CH₃OH); IR (KBr) 3327,
 1658, 1586, 1548, 1501, 1416, 1341, 1238, 1126; ¹H NMR
- 20 (CD₃OD) δ 7.22 (2H, s), 5.21 (1H, m), 5.00 (1H, m), 4.56, 4.49 (2H, 2m), 4.25 (1H, m), 3.88 (6H, s), 3.80 (3H, s), 3.55-3.43 (1H, m), 3.12 (1H, m), 2.71-1.70 (9H, m). Anal. Calcd for C₂₄H₃₀N₄O₁₀•2H₂O: C, 50.52; H, 6.01; N, 9.82. Found: C, 50.49; H, 6.05; N, 9.68. MS (ES⁺,
- 25 m/z) 533 (M+ 1, 1009).

[3S(1S,9S)] 3-[6,10-Dioxo-9-(naphth-1-oylamino)1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4cxobutanoic acid (408), was prepared by a similar
30 method to that described for compound 214e to afford a

white solid (73%): mp. 157-165°C (not sharp); $\left[\alpha\right]_0^{27} - 140.5^\circ$ (c 0.1, CH₃OH); IR (KBr) 3325, 1658, 1531, 1420, 1278, 1257; 1 H NMR (CD₃OD) δ 8.33-8.26 (1H, m), 8.01-7.78 (2H, m), 7.71 (1H, d, J = 6.0), 7.59-7.52 (3H, m), 5.27 (1H, m), 5.12-5.03 (1H, m), 4.55 (2H, m), 4.25 (1H, m), 3.64-3.43 (1H, m), 3.24-3.12 (1H, m), 2.80-1.67 (9H, m). Anal. Calcd for $C_25H_26N_4O_7*2H_2O$: C, 56.60; H, 5.70; N, 10.56. Found: C, 56.70; H, 5.80; N, 10.33. MS (ES^4 , m/z), 493 (M^4 - 1, 100%).

- 10 [3S(1S,9S)] 3-[6,10-Dioxo-4-(hydroxy-3,5dimethylbenzoyl)amino-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4oxobutanoic acid (214w), was prepared by a similar
 method as compound 214e to afford 210mg (62%) of a
- 15 white solid: mp. >260°C; $\left(\alpha\right)_{D}^{20}$ -93° (c 0.20, MeOH); IR (KBr) 3401, 2948, 1651, 1604, 1559, 1486, 1421, 1325, 1276, 1210; ^{1}H NMR (D₆-DMSO) δ 9.39 (1H, bs), 8.29 (1H, d, J = 5.9), 7.55 (2H, s), 6.64 (1H, d, J = 6.1), 5.79 (1H, s), 5.25-5.21 (1H, m), 1.90-1.82 (1H, m), 4.41-
- 20 3.69 (2H, m), 3.47-3.20 (3H, m), 2.97-2.91 (1H, m), 2.23 (6H, s), 2.25-1.60 (7H, m).

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213y R= Bn

[1S,9S(2RS,3S)] N-(2-Ethoxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)-1,2,3,4,7,8,9,10-

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octahydro-6-H-pyridazino[1,2-a][1,2]diazepine-1carboxamide (550q), was synthesized via methods used to prepare 213e to afford 550q.

[1S,95(2RS,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-35 yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)1,2,3,4,7,8,9,10-octahydro-6-Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (213y),
was synthesized via methods used to prepare 213e to
afford 213v.

10 [1s,9s(2s,3s)] N-(2-Phenethoxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]
diazepine-1-carboxamide, (412a) was synthesized via
methods used to prepare 550q using 513a-1 to afford
15 412a.

[15,95(2R,35)] N-(2-Phenethoxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]
diazepine-1-carboxamide, (412b) was synthesized via

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methods used to prepare 550q using 513a-2 to afford 412b.

- [1S,9S(2S,3S)] N-(2-Cyclopentoxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)-
- 5 1,2,3,4,7,8,9,10-octahydro-6-Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide, (412c) was synthesized via methods used to prepare 550q using 513b-1 to afford 412c.
- [1S,9S(2R,3S)] N-(2-Cyclopentoxy-5-oxotetrahydrofuran10 3-yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)1,2,3,4,7,8,9,10-octahydro-6-Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide, (412d)
 was synthesized via methods used to prepare 550q using
 513b-2 to afford 412d: ¹H NMR (CDCl₃) & 9.5 (1H, d),
- 15 8.9 (1H, d), 8.5 (1H, d), 7.9-7.8 (2H, m), 7.8-7.65 (2H, m), 6.55 (1H, d), 5.55 (1H, d), 5.25-5.1 (2H, m), 4.75-4.65 (1H, m), 4.65-4.6 (1H, m), 4.4-4.3 (1H, m), 3.25-3.15 (1H, m), 3.15-3.05 (1H, m), 2.95-2.8 (2H, m), 2.55-2.4 (2H, m), 2.15-1.5 (14H, m).
- 20 [18,98(28,38)] N-(2-Ethoxy-5-oxotetrahydrofuran-3-y1)6,10-dioxo-9-(isoquinolin-1-oylamino)-1,2,3,4,7,8,9,10octahydro-6-H-pyridazino[1,2-a][1,2] diazepine-1carboxamide, (412e) was synthesized via methods used to
 prepare 550q using 513f-1 to afford 412e.
- 25 [1s,9s(2R,3s)] N-(2-Ethoxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)-1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2] diazepine-1-

carboxamide, (412f) was synthesized via methods used to prepare 550q using 513f-2 to afford 412f.

Compounds 410 and 412 were prepared via methods used to prepare 605 from 604.

5 502y, 502z

410, 412

compound	R ¹
5 02 y, 410	S)
502z, 412	

[3S(1S,9S)] 3-[(6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-10 6H-pyridazino[1,2-a][1,2]diazepine-9-(thiophene-3-yl-carbonylamino)-1-carboxamido]-4-oxobutanoic acid (410), was purified by flash chromatography (5-254 methanol incichloromethane) to give 296mg (94%) of a colourless solid: mp. 90-200°C; IR (KBr) 3338, 3096, 2950, 1787, 15 1726, 1657, 1546, 1420, 1279, 1258, 1125, 1092, 984,

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933; 1 H NMR (CD₃OD) δ 8.41 (1H, d), 8.13 (1H, d), 7.54-7.41 (3H, m), 7.20 (1H, d), 5.19-5.11 (1H, m), 4.54-4.30 (1H, m), 3.27 (1H, m), 3.18-3.03 (1H, m), 2.81-2.64 (2H, m), 2.56-1.59 (7H, m). Anal. Calcd for 5 C₁₉H₂P₃V₄O₇S+2.5H₂O: C, 46.05; H, 5.49; N, 11.31. Found: C, 46.36; H, 5.25; N, 11.10. MS (ES⁺) 449 (M - 1, 80%), 113 (100). Accurate mass calculated for C₁H₂P₃V₄O₇S (MH⁺): 451.1287. Found: 451.1295.

[3S(1S,9S)] 3-[6,10-Dioxo-9-(isoquinolin-1-oylamino) 10 1,2,3,4,7,8,9,10-octahydro-6H pyridazıno[1,2-a][1,2]diazepine-1-carboxamido]-4 oxobutanoic acid (412) was prepared by a similar method
 to that described for compound 605 to afford a white
 glassy solid (69%): mp. 138-141°C; [α]_D²³ -105.5° (c
15 0.5, CH₂Cl₂); IR (KBr) 3375, 1787, 1659, 1515, 1421,
 1278, 1256; ¹H NMR (CDCl₃) δ 9.32 (1H, m), 8.79 (1H, m),
 8.47 (1H, m), 7.86-7.64 (4H, m), 5.31, 5.18, 4.59, 4.37
 (4 or 5H, m), 3.55-2.76, 2.49-2.39, 2.05, 1.65 (11H,
 4m). Anal. Calcd for C₂₄H₂₅N₅C₇+1.5H₂O: C, 55.17; H,

20 5.40; N, 13.40. Found: C, 54.87; H, 5.22; N, 13.15.

 $MS (ES^+, m/z) 494 (M^+ - 1, 100\%)$

[3s(1s,9s)] t-Butyl 3-[6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-9-(thiophene-3-yl)-6H-pyridazino[1,2-a][1,2]diazepine-carbonylamino)-125 carboxamido]-4-oxobutanoate semicarbazone (502y), was synthesized via methods used to prepare 604 from 603 to afford a pale cream powder: mp. 120-180°C; [α]_D²³ - 109° (c 0.18, CH₂Cl₂); IR (KBr) 3478, 3327, 1670, 1562, 1543, 1421, 1279, 1257, 1155; ¹H NNR (CDCl₃, CD₃OD) δ
30 8.04 (1H, m), 7.49 (1H, m), 7.38 (1H, m), 7.17 (1H, m).

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[3S(1S,9S)] t-Butyl 3-[6,10-dioxo-9-(isoquinolin-1oylamino)-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4oxobutanoate semicarbazone (502z), was prepared by a
similar method to that described for compound 604 to
afford a pale yellow solid (90%): mp. 142-145°C; [α]_D²⁴
15 -136.5° (c 0.06, CH₂Cl₂); ¹H NMR (CDCl₃) δ 9.51-9.46 (Hi,
m), 9.11 (HH, s), 8.83 (HH, d, J = 7.8), 8.53 (HH, d, J
= 5.5), 7.89-7.83 (2H, m), 7.77-7.65 (2H, m), 7.55 (HH,
d, J = 7.2), 7.18 (HH, d, J = 2.7), 5.26-5.12 (2H, m),
4.87 (HH, m), 4.59 (HH, m), 3.25-3.12 (2H, m), 2.9520 2.76 (2H, m), 2.59-2.38, 2.18-1.94, 1.70 (5H, 3m), 1.44

(9H, s).



compound	R ⁴	R ¹
415a	(II)	\n^()

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compound	R ⁴	R ¹
415b	an i	,0
415c	an i	~ ()
214w-1	CH ₃	,000
214w-2	CH ₃	,°`()
214w-3	CH ₃	
214w-4	HO CH,	,~~(<u>)</u>
214w-5	CH ₃	,~\\[\bar{\bar{\bar{\bar{\bar{\bar{\bar{
214w-6	CH ₉	,oC)
214w-7	CH ₃	,eQ
412g		,°``()

5

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compound	R ⁴	R ¹
412h		~0

[15,98(25,35)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(methylenedioxybenzoylamino)1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]
diazepine-1-carboxamide, (415a) was synthesized via
methods used to prepare 550g to afford 415a.

[1s,9s(2Rs,3s)] N-(2-Ethoxy-5-oxotetrahydrofuran-3-y1)6,10-dioxo-9-(methylenedioxy benzoylamino)1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]
10 diazepine-1-carboxamide, (415b) was synthesized via
methods used to prepare 550q to afford 415b.

[15,95(2R,35)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3yl)-6,10-dioxo-9-(methylenedioxy benzoylamino)1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]
diazepine-1-carboxamide, (415c) was synthesized via
methods used to prepare 550q to afford 415c.

[1s,9s(2Rs,3s)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-y1)-6,10-dioxo-9-(3,5-dimethyl-4-hydroxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-6-H-

20 pyridazino[1,2-a][1,2]diazepine-1-carboxamide, (214w :)
 was synthesized via methods used to prepare 550q to
 afford 214w-1.

[1S,9S(2R,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(3,5-dimethyl-4-hydroxybenzoylamino)-

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1,2,3,4,7,8,9,10-octahydro-6-H-

10 afford 214w-3.

pyridazino[1,2-a][1,2]diazepine-1-carboxamide, (214w-2)
was synthesized via methods used to prepare 550q to
afford 214w-2.

- [15,95(25,35)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(3,5-dimethyl-4-hydroxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide, (214w-3) was synthesized via methods used to prepare 550q to
 - $\label{eq:continuous} \begin{tabular}{ll} $\{1.5,9.5(2.R,3.5)\}$ &$N-(2-Phenethoxy-5-oxotetrahydrofuran-3-y1)-6,10-dioxo-9-(3,5-dimethyl-4-hydroxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-6-H-\\ \end{tabular}$
- pyridazino[1,2-a][1,2]diazepine-1-carboxamide, (214w-4)
 15 was synthesized via methods used to prepare 550q to
- afford 214w-4.
- [15,95(25,35)] N-(2-Phenethoxy-5-oxotetrahydrofuran-3-y1)-6,10-dioxo-9-(3,5-dimethyl-4-hydroxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-6-H-
- 20 pyridazino[1,2-a][1,2]diazepine-1-carboxamide, (214w-5)
 was synthesized via methods used to prepare 550q to
 afford 214w-5.
 - [1s,9s(2R,3s)] N-(2-Cyclopentoxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(3,5-dimethyl-4-
- 25 hydroxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-6-Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide, (214w-6)
 was synthesized via methods used to prepare 550q to
 afford 214w-6

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[1S,9S(2S,3S)] N-(2-Cyclopentoxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(3,5-dimethyl-4hydroxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-6-Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide, (214w-7) 5 was synthesized via methods used to prepare 550q to afford 214w-7.

[1S,9S(2R,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]
10 diazepine-1-carboxamide, (412g) was synthesized via

[1S,9S(2S,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3yl)-6,10-dioxo-9-(isoquinolin-1-oylamino)1,2,3,4,7,8,9,10-octahydro-6-H-pyridazino[1,2-a][1,2]
diazepine-1-carboxamide, (412h) was synthesized via
methods used to prepare 550g to afford 412h.

methods used to prepare 550g to afford 412g.

[3s(1s,9s)]3-(9-(4,5-Methylenedioxybenzoyl)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-420 oxobutanoic acid (415), was synthesized by the method used to prepare 2002 from 2001 to afford 415.

used to prepare 2002 from 2001 to afford 214w.

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[3S(1S,9S)]3-(9-(3,5-Dichloro-4-hydroxybenzoy1)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-45 oxobutanoic acid (214w), was synthesized by the method

2100k-o

compound	R
2100k	,°~~
21001	³ 0-⟨○
2100m	,°-(C)
2100n	ht. 0
21000	~~~

10

- 590 -

[1S,9S(2RS,3S)] 9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-N-(2-phenethyloxy-5-oxotetrahydrofuran-3-yl)-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (2100k),

- 5 was prepared by a similar method as compound 213e to afford a mixture of diastereoisomers (75/25) as a white solid (258mg, 83%): mp. 101°C ; $\left[\alpha\right]_{D}^{25}$ -96° (c 0.2, $\text{CH}_{2}\text{Cl}_{2}$); IR (KBr) 3328, 2935, 2978, 1732, 1669, 1603, 1483, 1450, 1414, 1237, 1155, 1082, 989, 755; $^{1}\text{H}_{NMR}$
- 10 (CDC1₃) δ 7.84-7.80 (2H, m), 7.54-7.17 (8H, m), 7.06-6.99 (1H, m), 6.25 (1H, d, J = 7.9H), 5.41 (0.75H, d, J = 5.4H), 5.31 (0.25H, bs), 5.23-5.09 (1H, m), 4.93-4.87 (1H, m), 4.68-4.51 (2H, m), 4.40-4.33 (0.25H, m), 4.24-4.14 (0.75H, m), 3.95-3.70 (1H, m), 3.30-3.13 (1H, m),
- 15 3.14-2.78 (5H, m), 2.47-2.21 (2H, m), 2.05-1.50 (5H, m). Anal. Calcd for $C_{29}H_{32}N_4O_7\cdot 0.5H_2O$: C, 62.47; H, 5.97; N, 10.05. Found: C, 62.17; H, 5.83; N, 9.97. MS (ES $^+$) 549.

[1S,9S(2RS,3S)] 9-Benzamido-N-(2-cyclopentyloxy-5-oxo-

- 20 tetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide (21001), was prepared by a similar method as 213e, (74%) as a colourless solid: mp. 172-80°C; [\alpha]_p^{23} -91.5° (c 0.1, CH₂Cl₂); IR (KBr) 3290, 1792,
- 25 1677, 1657, 1642, 1544, 1425, 1280, 1259, 1124, 977; ²H
 NMR (CDCl₃) δ7.80 (2H, m), 7.46 (3.5H, m), 7.00 (1H, d, J = 6.7), 6.48 (0.5H, d, J = 7.9), 5.55 (0.5H, d, J = 5.3), 5.19 (2H, s + m), 4.93 (0.5H, m), 4.62 (1.5H, m), 4.34 (1H, m), 4.18 (0.5H, m), 3.28-2.70 (4H, m), 2.49-
- 30 2.29 (2H, m), 205-1.48 (15H, m).

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[1s,9s(2R,3s)] 9-Benzamido-6,10-dioxo-N-[2-(2-indanyloxy)-5-oxo-tetrahydrofuran-3-yl]1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamide (2100m),

- 5 was prepared by a similar method as 213e, (76%) as a colourless solid: mp. ~140°C, remelts 187-9°C; $\left[\alpha\right]_{\rm D}^{23}-96.9$ ° (c 0.11, CH₂Cl₂); IR (KBr) 3507, 3308, 3251, 1772, 1660, 1641, 1566, 1545, 1457, 1424, 1346, 1326, 1302, 1275, 1258, 1136, 1085, 1018, 981; 1 H NMR (CDCl₃)
- 10 δ 7.78 (2H, m), 7.53 (3H, m), 7.19 (4H, m), 6.91 (1H, d, J = 7.4), 6.27 (1H, d, J = 7.6), 5.66 (1H, d, J = 5.3), 5.10 (1H, m), 4.96 (1H, m), 4.75 (2H, m), 4.52 (1H, m), 3.08 (3H, m), 3.03-2.71 (5H, m), 2.48-2.31 (2H, m), 1.90-1.40 (4H, m), 1.22 (1H, m).
- 15 [15,95(25,35)] 9-Benzoylamino-N-(2-benzyloxy-5oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1carboxamide (2100n), was prepared by a similar method
 to that described for compound 213e to afford a white
- 20 glassy solid (76%): mp. $112-5^{\circ}\text{C}$; $\{\alpha\}_0^{23} -62.0^{\circ}$ (c 0.1, CH₂Cl₂); IR (KBr) 3305, 1789, 1677, 1665, 1535, 1422, 1279, 1256, 1119, 942, 700; ^{1}H NMR (CDCl₃) δ 7.84 (2H, m), 7.58-7.27 (9H, m), 6.99 (1H, d, \mathcal{J} = 7.8), 5.23 (1H, s), 5.23-5.11 (1H, m), 4.89 (1H, m), 4.76 (1H, d, \mathcal{J} =
- 25 11.3), 4.55 (1H, d, J = 11.4), 4.56-4.43 (2H, m), 3.30-2.96, 2.81-2.69, 2.46-2.37, 2.16-1.66 (10H, 4m), 2.27 (1H, d, J = 17.8). Anal. Calcd for C₂₈H₃₀N₄O₇·0.5H₂O: C, 61.87; H, 5.75; N, 10.32. Found: C, 61.88; H, 5.70; N, 10.33. MS (ES⁺, m/z) 535 (M⁺ + 1, 100%).

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[1S,9S(2R,3S)] 9-Benzoylamino-N-(2-benzyloxy-5oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1carboxamide (2100o), (containing about 7% of (2S)), was 5 prepared by a similar method to that described for compound 213e to afford a white glassy solid (81%): mp. 115-7°C; $[\alpha]_n^{23}$ -121.8° (c 0.11, CH₂Cl₂); IR (KBr) 3326, 1792, 1659, 1535, 1421, 1278, 1257, 1124, 978; ¹H NMR (CDCl₃) δ7.82 (2H, m), 7.58-7.24 (8H, m), 6.90 (1H, 10 d, J = 7.3), 6.49 (1H, d, J = 7.7), 5.57 (1H, d, J = 7.7) 5.5), 5.11 (2H, m), 4.91 (1H, d, J = 11.4), 4.57 (1H, d, J = 11.1), 4.81-4.68 (1H, m), 4.65-4.54 (1H, m), 3.18-2.71 2.52-2.30, 2.05-1.62 (11H, 3m). Anal. Calcd for C28H30N4O7 • 0.5H2O: C, 61.87; H, 5.75; N, 10.32. 15 Found: C, 61.70; H, 5.71; N, 10.15. MS (ES+, m/z) 535 $(M^+ + 1, 94.3\%), 557 (100\%).$

550n

[1S,9S(2RS,3S)] 9-(3-Acetamido)benzoylamino-6,10-dioxo-N-(2-ethoxy-5-oxo-tetrahydrofuran-3-yl)-

20 1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (550n),
was prepared by a similar method as compound 213e to

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afford a mixture of diastereoisomers (65/35) as a tan powder (390mg, 28%): mp. 139-145°C; [α]_D²³ -104° (c 0.2, MeOH); IR (KBr) 3318, 2405, 2369, 1792, 1660, 1591, 1549, 1484, 1422, 1257, 1117; 1 H NMR (D_6 -DMSO) δ 5 10.1 (1H, s), 8.80 (0.65H, d, J = 6.6), 8.58 (0.35H, d, J = 6.6), 8.59 (1H, d, J = 7.0), 8.06 (1H, bs), 7.83-7.79 (1H, m), 7.61-7.57 (1H, m), 7.47-7.39 (1H, m), 5.61 (0.35H, d, J = 5.0), 5.37 (0.65H, bs), 5.17-5.14 (0.35H, m), 5.08-5.06 (0.65H, m), 4.92-4.86 (1H, m), 10.467-4.61 (0.35H, m), 4.47-4.41 (0.65H, m), 4.28-4.11 (1H, 2m), 3.80-3.59 (2H, m), 3.23-2.75 (3H, m), 2.61-1.48 (7H, m), 2.10 (3H, s), 1.25 and 1.17 (3H, 2t, J = 5.8). MS (ES*) 528.

5500

15 [1s,9s(2Rs,3s)] 6,10-Dioxo-N-(2-ethoxy-5-oxotetrahydrofuran-3-yl)-9-(2-indoloylamino)1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (550o),
was synthesized by a similar method as compound 213e to
20 afford a colourless solid (1.071g, 80%): mp. 155-70°C;
[α]_D²²-75.8° (c 0.26, CH₂Cl₂); IR (KBF) 3314, 2941,
1791, 1658, 1545, 1420, 1341, 1312, 1252, 1181, 1118,
939, 749; ¹H NMR (CDCl₃) δ 9.45 (0.5R, s), 9.34 (2.5H, s), 7.68-7.62 (1H, m), 7.49-7.39 (2H, m), 7.33-7.26

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(1H, m), 7.18-7.03 (3H, m), 5.49 (0.5H, d), 5.30 (0.5 H, s), 5.26-5.13 (1H, m), 4.90-4.83 (0.5H, m), 4.76-4.49 (1H, m), 4.42-4.35 (0.5H, m), 3.97-3.74 (1H, m), 3.72-3.53 (1H, m), 3.35-2.64 (4H, m), 2.50-2.37 (1H, 5 m), 2.20-1.82 (5H, m), 1.69-1.50 (2H, m), 1.30-1.19 (3H, m).

550p

[1s,9s(2Rs,3s)] 6,10-Dioxo-N-(2-ethoxy-5-oxotetrahydrofuran-3-yl)-9-(4-hydroxybenzoyl)amino10 1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamide (550p),
was prepared by a similar method as compound 213e to
afford a mixture of diastereoisomers as a white foam
(820mg, 47%): [α]_D²⁴ -75° (c 0.16, CH₂Cl₂); IR (KBr)
15 3401, 2937, 1791, 1657, 1609, 1539, 1505, 1423, 1277,
1177, 1118; ¹H NMR (CDCl₃)δ8.07-8.05 (1H, m), 7.67 (2H,
d, J = 7.9), 7.38-7.29 (2H, m), 6.80 (2H, d, J = 8.5),
5.49 (0.5H, d, J = 4.6), 5.23 (0.5H, bs), 5.24-5.20
(1H, m), 5.12-5.08 (1H, m), 4.68-4.29 (2H, m), 3.9220 3.45 (3H, m), 3.32-2.30 (2H, m), 2.80-1.56 (11H, m),
1.21 (3H, t, J = 7.0H).

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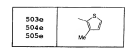
286, 505b-e

compound	R
503a 504a 286	
503b 504b 505b	Me Ph
503c 504c 505c	OPh
503d 504d 505d	OPh

5

10

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[3S,4R(1S,9S)] t-Butyl 3-(6,10-dioxo-9-

- 5 methanesulphonylamino-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4hydroxy-5-(1-naphthoyloxy)pentanoate (503a), was
 prepared from 212b and (35,4R) t-butyl (Nallyloxycarbonyl)-3-amino-4-hydroxy-5-(1-
- 10 naphthoyloxy)pentanoate by the method described for (213e) to afford 533mg (81%) of an off-white foam: $\left[\alpha\right]_D^{22} 81.4^\circ \text{ (c 0.5, CH}_2\text{Cl}_2); \text{ IR}(\text{KBr}) \text{ 3342, 2976, 1719,} \\ 1664, 1328, 1278, 1246, 1153, 1137. \\ \hline \\ 18.86 \text{ (1H, d, J = 8.4), 8.21 (1H, dd, J = 1.3, 7.3),}$
- 15 8.03 (1H, d, J = 8.1), 7.88 (1H, d, J = 8.6), 7.66-7.45 (3H, m), 7.23 (1H, d, J = 8.6), 5.96 (1H, d, J = 9.2), 5.30 (1H, m), 4.59-4.33 (5H, m), 4.24 (1H, m), 3.96 (1H, brd), 3.29 (1H, m), 2.95 (1H, m), 2.93 (3H, s), 2.69-2.50 (3H, m), 2.36 (1H, m), 1.96 (4H, m), 1.62
- 20 (1H, m), 1.41 (9H, s). Anal. Calcd for $C_{31}H_{40}N_4O_{10}S \cdot 0.25H_{2}O$: C, 55.97; H, 6.14; N, 8.42. Found: C, 55.90; H, 6.11; N, 8.23. M.S. (ES^T) 683 (M+Na, 100%), 661 (M+1,39), 605 (78).

[3S(1S,9S)] t-Butyl 3-(6,10-dioxo-9-

25 methanesulphonylamino-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(1naphthoyloxy)-4-oxopentanoate (504a), was synthesized
from 503a via method used to prepare 216e from 215e to
afford 446mg (91%) of a colourless foam: [\alpha]n²¹ -111.6°

10 Calcd for $C_{31}H_{38}N_4O_{10}S \cdot 0.25H_2O$. C, 56.14; H, 5.85; N, 8.45. Found: C, 56.11; H, 5.83; N, 8.29. M.S. (ES*) 657 (M-1, 100%).

[3S(1S,9S)] 3-(6,10-Dioxo-9-methanesulphonylamino-1,2,3,4,7,8,9,10-octahydro-6H-

- pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(1-naphthoyloxy)-4-oxopentanoic acid (286), was prepared from 504a by the method described for 217 to afford 356mg (93%) of a white powder: mp 120-123°C; $\left[\alpha\right]_{D}^{23}$ 121° (c 0.194, CH₂Cl₂); IR (KBr) 3314, 2937, 1722,
- 20 1663, 1412, 1328, 1278, 1245, 1195, 1132.
 ¹H NMR (d6-DMSO) δ12.63 (1H, brs), 8.94 (1H, d, J = 7.4), 8.78 (1H, d, J = 8.6), 8.26 (2H, m), 8.11 (1H, d, J = 8.0), 7.77-7.62 (4H, m), 5.28 (2H, s), 5.21 (1H, m), 4.82 (1H, m), 4.44-4.29 (2H, m), 3.31 (1H, m), 2.98 (3H, s), 2.98-
- 25 2.86 (2H, m), 2.72 (1H, dd, J = 7.3, 16.9), 2.40 (1H, m), 2.24-1.84 (4H, m), 1.69 (2H, m). Anal. Calcd for $C_{27}H_{30}N_4O_{10}S\cdot H_2O: C, 52.25; H, 5.20; N, 9.03. Found: C, 52.11; H, 4.97; N, 8.89. M.S. (ES +) 601 (M-1, 100%).$
- 30 [35,4RS(1S,9S)] t-Butyl 3-[6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2]diazepine-9-

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15 [3s(1s,9s)] t-Butyl 3-[6,10-dioxo-9(methanesulphonylamino)-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-5-(5methyl-3-phenylisoxazoyloxy)-4-oxopentanoate (504b),
was synthesized by a similar method as compound 216b to
20 afford a colourless powder (601mg, 93%): mp. 75-115°C;
[α]₂²³ -104° (c 0.26, CH₂Cl₂); IR (KBr) 3324, 2977,
2935, 1730, 1670, 1525, 1452, 1422, 1369, 1317, 1276,
1256, 1222, 1155, 1107, 990, 766; ¹H NMR (CDCl₃) δ 7.687.61 (2H, m), 7.47-7.38 (3H, m), 7.32-7.24 (1H, m),
25 5.56 (1H, d), 5.36-5.24 (1H, m), 5.04 (1H, d), 4.88

(1H, d), 4.86-4.77 (1H, m), 4.64-4.39 (2H, m), 3.32-3.17 (1H, m), 2.97-2.85 (1H, m), 2.93 (3H, s), 2.76 (3H, s), 2.80-2.71 (1H, m), 2.65-2.49 (1H, m), 2.41-2.30 (1H, m), 2.12-1.61 (6H, m), 1.42 (9H, s). Anal.

30 Calcd for C₃₁H₃₉N₅O₁₁S·H₂O: C, 52.61; H, 5.84; N, 9.90; S, 4.53. Found: C, 52.94; H, 5.69; N, 9.72; S, 4.51.

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MS (ES^+) 712 (31%), 707 (100), 690 $(M^+ + 1, 41)$, 634 (55).

[3S(1S,9S)] 3-[6,10-Dioxo-9-(methanesulphonylamino)-1,2,3,4,7,8,9,10-octahydro-6H-

- 5 pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-5-(5-methyl-3-phenylisoxazoyloxy)-4-oxopentanoic acid (505b), was synthesized by a similar method as compound 217 to afford a colourless powder (499mg, 96%): mp. 95-145°C; [α]_D²² -137° (c 0.12, MeOH); IR (KBr) 3323,
- 10 2936, 1732, 1665, 1529, 1452, 1421, 1312, 1275, 1256, 1221, 1183, 1153, 1135, 1101, 990; ^{1}H NMR (CD₃OD) δ 7.67-7.56 (2H, m), 7.49-7.38 (4H, m), 5.23-5.12 (1H, m), 5.02 (1H, d), 4.79-4.73 (1H, m), 4.52-4.34 (3H, m), 3.48-3.25 (2H, m), 3.03-2.85 (2H, m), 2.94 (3H, s),
- 15 2.74 (3H, s), 2.79-2.66 (1H, m), 2.52-2.38 (1H, m),
 2.29-2.14 (1H, m), 2.04-1.70 (4H, m). Anal. Calcd for $C_{27}H_{31}N_{5}O_{11}S \cdot H_{2}O$: C, 49.77; H, 5.18; N, 10.75; S, 4.92.
 Found: C, 49.83; H, 5.01; N, 10.27; S, 4.84. MS (ES⁺)
 746 (42%), 632 (M 1, 100), 386 (60). Accurate mass
- 20 calculated for $C_{27}H_{32}N_5O_{11}S$ (MH $^+$): 634.1819. Found: 634.1807

[3S,4RS(1S,9S)] t-Butyl 3-[6,10-dioxo-9-(methanesulphonylamino)-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-

- 25 hydroxy-5-(2-phenoxybenzoyloxy)pentanoate (503c), was synthesized by a similar method as compound 213e to afford a colourless solid (446mg, 84%): IR (KBr) 3345, 2976, 2935, 1727, 1664, 1603, 1535, 1483, 1451, 1416, 1395, 1369, 1328, 1297, 1277, 1237, 1155, 1135, 1076,
- 3C 990, 755; ¹H NMR (CDCl₃) δ7.98-7.89 (1H, m), 7.55-7.45

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[3S(1S,9S)] t-Butvl 3-[6,10-dioxo-9-

(methanesulphonylamino) -1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-(2-phenoxybenzovloxy) pentanoate (504c), was

- 15 synthesized by a similar method as compound 216e to afford a colourless powder: mp. 85-100°C; $\left[\alpha\right]_0^{22}$ -91.3° (c 0.52, CH₂Cl₂); IR (KBr) 3328, 2978, 2935, 1732, 1669, 1603, 1524, 1483, 1450, 1396, 1369, 1296, 1276, 1237, 1155, 1132, 1082, 989, 755; 1 H NMR (CDCl₃) δ 8.03-
- 20 7.98 (1H, m), 7.52-7.44 (1H, m), 7.37-7.07 (5H, m), 7.01-6.92 (3H, m), 5.52 (1H, d), 5.26-5.20 (1H, m), 5.06-4.84 (3H, m), 4.64-4.39 (2H, m), 3.32-3.14 (1H, m), 2.99-2.88 (1H, m), 2.94 (3H, s), 2.65-2.45 (2H, m), 2.39-2.29 (1H, m), 2.12-1.58 (6H, m), 1.40 (9H, s).
- 25 Anal. Calcd for $C_{33}H_{40}N_4O_{11}S$: C, 56.56; H, 5.75; N, 8.00; S, 4.58. Found: C, 56.37; H, 5.84; N, 7.69; S, 4.37. MS (ES^+) 723 (30%), 718 (100), 701 IM^+ + 1, 23), 645 (59).

[3S(1S,9S)] 3-[6,10-Dioxo-9-(methanesulphonylamino)30 1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-

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(2-phenoxybenzoyloxy)pentanoic acid (505c), was synthesized by a similar method as compound 217 to afford a colourless foam (252mg, 72%): mp. 90-125°C; $[\alpha]_D^{23}$ -133° (c 0.11, MeOH); IR (KBr) 3314, 2938, 5 1792, 1734, 1663, 1604, 1535, 1483, 1448, 1415, 1250, 1132, 756; 1 H NMR (D_6 -DMSO) & 8.61-8.76 (1H, m), 7.92 (1H, d), 7.68-7.54 (2H, m), 7.41-7.25 (3H, m), 7.16-6.91 (4H, m), 5.13-4.98 (2H, m), 4.72-4.63 (1H, m), 4.37-4.21 (2H, m), 2.92 (3H, s), 2.90-2.60 (3H, m), 10 2.35-2.26 (1H, m), 2.17-2.05 (2H, m), 1.99-1.80 (2H,

m), 1.61-1.50 (1H, m).Anal. Calcd for $C_{29}H_{32}N_4O_{11}S \cdot 0.5H_2O: C, 53.29; H, 5.09; N, 8.57; S, \\ 4.90. Found: C, 53.57; H, 5.18; N, 8.32; S, 4.75. MS \\ (ES^+) 643 (M-1, 100%).$

- 7.00 (2H, d), 5.93-5.80 (1H, m), 5.36-5.30 (1H, m), 25 4.63-4.24 (5H, m), 4.15-4.09 (1H, m), 5.37-3.22 (1H, m), 2.98-2.74 (1H, m), 2.94 (3H, s), 2.70-2.47 (3H, m), 2.40-2.30 (1H, m), 2.15-1.60 (5H, m), 1.42 (9H, s). Anal. Calcd for C₃₃H₄₂N₄O₁₁S·H₂O: C, 54.99; H, 6.15; N, 7.77; S, 4.45. Found: C, 54.60; H, 5.88; N, 7.49; S,

(CDCl₃) 87.77 (1H, d), 7.67 (1H, m), 7.45-7.10 (6H, m),

30 4.50. MS (ES⁺) 725 (19 \S), 720 (91), 703 (M⁺ + 1, 74), 647 (76), 629 (100), 433 (78).

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[3S(1S,9S)] t-Butyl 3-[6,10-dioxo-9-(methanesulphonylamino)-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-(3-phenoxybenzoyloxy) pentanoate (504d), was

- 5 synthesized by a similar method as compound 216e to afford a colourless powder (466mg, 85%): mp. 75-100°C; [α]_D²² -99.3° (c 0.60, CH₂Cl₂); IR (KBr) 3335, 2978, 2937, 1728, 1669, 1584, 1525, 1487, 1444, 1416, 1369, 1328, 1272, 1227, 1188, 1155, 989, 754; ¹H NMR (CDCl₃) δ 10 7.82-7.77 (1H, m), 7.66-7.65 (1H, m), 7.46-7.32 (4H, m), 7.26-7.10 (2H, m), 7.04-6.98 (2H, m), 5.68 (1H, d), 5.37-5.31 (1H, m), 5.11 (1H, d), 5.02-4.88 (2H, m), 4.66-4.42 (2H, m), 3.3*-3.17 (1H, m), 2.98-2.89 (1H, m), 2.96 (3H, s), 2.84-2.78 (1H, m), 2.72-2.47 (1H, m),
- 15 2.42-2.32 (1H, m), 2.14-1.58 (6H, m), 1.43 (9H. s). Anal. Calcd for $C_{33}H_{40}N_{4}O_{11}S$: C, 56.56; H, 5.75; N, 8.00. Found: C, 56.36; H, 5.82; N, 7.71. MS (ES[†]) 723 (56%), 718 (90), 701 (M^{\dagger} + 1, 36), 645 (100).

[3s(1s,9s)] 3-[6,10-Dioxo-9-(methanesulphonylamino)-20 1.2.3.4.7.8.9,10-octahydro-6H-

- pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-(3-phenoxybenzoyloxy)pentanoic acid (505d), was synthesized by a similar method as compound 217 to afford a colourless foam (353mg, 73%): mp. 80-115°C; 25 [α]_D²³ -138° (c 0.11, MeOH); IR (KBr) 3327, 2937, 1728, 1666, 1584, 1529, 1487, 1443, 1413, 1328, 1273, 1227, 1189, 1155, 1134, 989, 754; ¹H NMF (D₆-DMSO) δ 8.92
- 1189, 1155, 1134, 989, 754;

 ¹H NMF (D₆-DMSO) δ 8.82 (1H, d), 7.76-7.72 (1H, m), 7.61-7.53 (2H, m), 7.48-7.32 (4H, m), 7.24-7.17 (1H, m), 7.11-7.06 (2H, m), 30 5.14-5.06 (3H, m), 4.73-4.64 (1H, m), 4.38-4.24 (2H, m)
- 30 5.14-5.06 (3H, m), 4.73-4.64 (1H, m), 4.38-4.24 (2H, m), 2.92 (3H, s), 2.89-2.61 (3H, m), 2.38-2.27 (1H, m),

2.19-2.06 (2H, m), 2.02-1.79 (3H, m), 1.63-1.52 (1H, m). Anal. Calcd for $C_{29}H_{32}N_{4}O_{11}S \cdot 0.5H_{2}O$: C, 53.29; H, 5.09; N, 8.57; S, 4.90. Found: C, 53.24; H, 5.14; N, 8.34; S, 4.86. MS (ES⁴) 643 (M - 1, 100%), 385 (62).

- 5 [3s,4R(1s,9s)] t-Butyl 5-(3-chlorothien-2-oyloxy)-3(6,10-dioxo-9-methanesulphonylamino-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1carboxamido)-4-hydroxypentanoate (503e), was prepared
 by a similar method to that described for compound
- 10 213e, to afford an off white solid (70%): mp. 100- 103° C; $\{\alpha\}_{D}^{25}$ -84.0° (c 0.05, CH₂Cl₂); IR (KBr) 3459- 3359, 1722, 1664, 1514, 1368, 1326, 1278, 1247, 1155; 1 H NMR (CDCl₃) δ 7.52 (1H, m), 7.06-6.99 (2H, m), 5.69 (1H, d, J = 9.0), 5.23 (1H, m), 4.61-4.16 (6H, m),
- 15 3.36-3.19 (1H, m), 2.96 (3H, s), 2.67-2.49, 2.42-2.32, 2.06-1.89, 1.69 (10H, 4m), 1.43 (9H, s).

[3s(1s,9s)] t-Butyl 5-(3-chlorothien-2-oyloxy)-3-(6,10-dioxo-9-methanesulphonylamino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-

- 20 carboxamido)-4-oxopentanoate (504e), was prepared by a similar method to that described for compound 216e, to afford a white solid (98%): mp. 91-98°C; (α)_D²⁵ 112.5°C (c 0.06, CH₂Cl₂); IR (KBr) 3453-3364, 1727, 1668, 1513, 1420, 1368, 1245, 1155; ¹H NMR (CDCl₃) 87.54
- 25 (1H, d, J = 5.3), 7.18 (1H, d, J = 7.18), 7.05 (1H, a, J = 5.4), 5.42 (1H, d, J = 8.9), 5.25 (1H, m), 5.02 (2H, m), 4.96-4.87 (1H, m), 4.65-4.42 (2H, m), 3.34-3.17 (1H, m), 2.97-2.93 (1H, m), 2.97 (3H, s), 2.87-2.78, 2.73-2.50, 2.38-2.32, 2.13-1.88, 1.69-1.60 (9H,
- 30 5m), 1.44 (9H, s).

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[3S(1S,9S)] 5-(3-Chlorothien-2-ovloxy)-3-(6,10-dioxo-9methanesulphonylamino-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4oxopentanoic acid (505e). A solution of 217 (0.33g, 5 0.51mmol) in dry dichloromethane (3ml) was cooled (ice/water) with protection from moisture. Trifluoroacetic acid (2ml) was added with stirring. The solution was kept at room temperature for 2h after removal of the cooling bath, then concentrated in 10 vacuo. The residue was evaporated three times from dichloromethane, triturated with diethyl ether and filtered. The solid was purified by flash chromatography (silica gel, 0-6% methanol in dichloromethane) to give the product as a white glassy 15 solid (0.296g, 98%): mp 110-122°C; $[\alpha]_D^{22}$ -163.5° (c 0.1, CH₃OH); IR (KBr) 3514-3337, 1726, 1664, 1513, 1420, 1245, 1152, 1134, 990; ¹H NMR (CD₃OD) δ7.79 (1H, d, J = 5.2), 7.12 (1H, d, J = 5.2), 5.20 (1H, m), 5.02-4.72 (2H, m, masked by H2O), 4.59-4.32 (3H, m), 3.48-20 3.29, 3.08-2.75, 2.50-2.41, 2.31-2.22, 2.08-1.89, 1.72-

1.63 (11H, 6m), 2.95 (3H, s).

506a-c,q 507a-c,q

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 compound
 R1

 506a
 PhC(0)

 507a
 PhC(0)

 506b
 MeS(0)2

 507c
 MeOC(0)

 506g
 CH3C(0)

 507a
 CH3C(0)

5

10 [3S(1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-diazo-4-oxopentanoate (506a). A solution of 212e (321mg, 0.929mmol) and (3S) t-butyl 3-amino-5-diazo-4-

- 15 oxopentanoate (198mg, 0.929mmol) in dichloromethane (3ml) was cooled to 0° and N,N-diisopropylethylamine (0.16ml, 1.86mmol) and [2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyl-uronium tetrafluoroborate (328mg, 1.02mmol) were added. The solution was stirred
- 20 overnight at room temperature, diluted with ethyl acetate and washed with 1M NaHSO₄ (x2), aqueous NaHCO₃ (x2), brine, dried over magnesium sulphate and evaporated. Chromatography on silica gel eluting with ethyl acetate gave 506a (425mg, 85%) as a colourless
- 25 foam: $\left[\alpha\right]_{D}^{23}$ -124.9° (c 0.2, CH₂Cl₂); IR (KBr) 3332, 2111, 1728, 1658, 1532, 1421, 1392, 1367, 1279, 1256, 1155; 1 H NMR (CDCl₃) δ 7.82 (2H, m), 7.49 (3H, m), 7.28 (1H, d, J = 9.3), 7.05 (1H, d, J = 7.3), 5.06 (1H, s), 5.18 (2H, m), 4.78 (1H, m), 4.62 (1H, m), 3.29 (1H, m), 30 3.08-2.79 (3H, m), 2.58 (1H, dd, J = 16.8, 5.6), 2.20-

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1.85 (4H, m), 1.70 (1H, m), 1.45 (9H, s). MS (ES^{+}) 539.58 (M - 1, 97.9%) 529.59 (100).

[3S(1S,9S)] t-Butyl 5-diazo-3-[6,10-dioxo-(9-methanesulphonamido)-1,2,3,4,7,8,9,10-octahydro-6H5 pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4 oxopentanoate (506b), was prepared by a similar method as compound 506a. 74% as yellow orange solid: mp. 75°C (decomp.); [α]_D²⁰ -92.0° (c 0.036, CH₂Cl₂); IR (KBr) 3438, 2904, 2113, 1728, 1669, 1523, 1368, 1328, 1155; 10

1H NMR (CDCl₃) δ 7.48 (1H, d, J = 8.1), 5.83-5.68 (1H, m,), 5.55-5.50 (1H, m), 5.43-5.14 (1H, m), 4.83-4.45 (3H, m), 3.40-3.19 (1H, m), 2.98 (3H, s), 2.92-2.30

[3S(1S,9S)] t-Butyl 5-diazo-3-[6,10-dioxo-(9-15 methoxycarbonyl)amino-1,2,3,4,7,8,9,10-octahydro-6H-

(4H, m), 2.24-1.70 (6H, m), 1.43 (9H, s).

[3S(1S,9S)] t-Butyl 3-(9-acetylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazıno[1,2-a][1,2]diazepine-1-carboxamido)-5-diazo-4-oxopentanoate (506g), was prepared by a similar

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method as compound 506a. 81%: $\left[\alpha\right]_D^{28}$ -146.7° (c 0.4, CH₂Cl₂); IR (KBr) 3438, 2904, 2113, 1728, 1669, 1523, 1368, 1328, 1155; 1 H NMR (CDCl₃) δ 7.32 (1H, d), 6.43 (1H, d), 5.50 (1H, s), 5.22 (1H, m), 4.94 (1H, m), 4.77 (1H, m), 4.60 (1H, m), 3.24 (1H, m), 3.03-2.52 (4H, m), 2.36 (1H, m), 2.10-1.64 (5H, m), 2.02 (3H, s), 1.45 (9H, s). Anal. Calcd for C₂₁H₂₀N₆O₇: C, 52.69; H, 6.32; N, 17.05. Found: C, 52.51; H, 6.27; N, 17.36. MS (ES⁵) 477 (M⁵ - 1, 100%).

- 10 [3S(1S,9S)] t-Butyl 5-bromo-3-(9-benzoylamino-6,10dioxo-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4oxopentanoate (507a). 506a (3.0g, 5.55mmol) in dry
 dichloromethane (40ml) was cooled to 0° and 30%
- 15 hydrobromic acid in acetic acid (1.1ml, 5.55mmol) was
 added dropwise over 4min. The mixture was stirred at
 0° for 9min and quenched with aqueous sodium
 bicarbonate. The product was extracted into ethyl
 acetate, washed with aqueous sodium bicarbonate, brine,
 20 dried (MgSO₄) and evaporated to give 2.97g (92%) of a
- colourless foam: [α]_D²³ -82.3° (c 0.23, CH₂Cl₂); IR (KBr) 3333, 1726, 1659, 1530, 1458, 1447, 1422, 1395, 1368, 1279, 1256, 1222, 1155, 728; ¹H NMR (CDCl₃; δ7.81 (2H, m), 7.50 (3H, m), 7.11 (1H, d, J = 8.0), 7.01 (1H,
- 25 d, J = 7.4), 5.20 (2H, m), 5.00 (1H, m), 4.06 (2H, s), 3.28 (1H, m), 3.20-2.70 (4H, m), 2.42 (1H, m), 2.10-1.85 (4H, m), 1.72 (1H, m), 1.44 (9H, s). Anal. Calcd for C₂₆H₃₃N₄O₇Br·C.7H₂O: C, 51.53; H, 5.72 N, 9.24. Found: C, 51.55; H, 5.52; N, 9.09. MS (ES[†]) 598. 593
- $30 (M^+ + 1)$.

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[3S(1S,9S)] t-Butyl 5-bromo-3-(6,10-dioxo-9-methanesulphonamido-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxopentanoate (507b), was prepared by a similar method 5 as compound 507a. (68%) as an orange foam: [α]_D²⁰ - 135° (c 0.053, CH₂Cl₂); IR (KBr) 3429, 2944, 2935, 1723, 1670, 1458, 1408, 1327, 1225, 1154, 991; ¹H MMR (CDCl₃) δ7.38 (1H, d, J = 8.2), 5.69 (1H, d, J = 9.3), 5,43-5.34 (1H, m), 5.07-4.97 (1H, m), 4.70-4.42 (2H, 10 m), 4.12 (2H, s), 3.35-3.17 (1H, mi), 3.10-2.69 (4H, mi), 2.98 (3H, s), 2.43-2.33 (1H, m), 2.15-1.65 (5H, m), 1.43 (9H, s). Anal. Calcd for C₂₀H₃₁BrN₄O₈S: C, 42.33; H, 5.51; N, 9.87. Found: C, 42.69; H, 5.52; N, 9.97.

[3S(1S,9S)] t-Butyl 5-bromo-3-(6,10-dioxo-9-

- 15 (methoxycarbonyl)amino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxopentanoate (507c), was prepared by a similar method as compound 507a to afford a pale yellow foam (320mg, 78\u00ea): [α]_D²⁰ -107° (c 0.2, CH₂Cl₂); IR (KBr) 3401,
 20 2956, 1726, 1670, 1528, 1452, 1415, 1395, 1368, 1276, 1251, 1155, 1064; ¹H NMR (CDCl₃) δ 7.07 (1H, d, J = 7.6), 5.47 (1H, d, J = 8.1), 5.21-5.16 (1H, m), 5.03-4.94 (1H, m), 4.75-4.56 (2H, m), 4.06 (2H, s), 3.69 (3H, s), 3.31-3.13 (1H, m), 3.03-2.92 (2H, m), 2.81-2.58 (2H, m), 2.41-2.31 (1H, m), 2.10-1.66 (5H, m), 1.44 (9H, s).
- [35(15,95)] t-Butyl 3-(9-acetylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-5-bromo-30 4-oxopentanoate (507g), was prepared by a similar

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compound	R
508a 284	CI
508b 285	Me

15 [3S(1S,9S)] t-Butyl 5-(2,6-dichlorobenzoyloxy)-3-[6,10-dioxo-9-(methoxycarbonyl)amino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxobutanoate (508a). To a solution of

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506c (547mg, 1mmol) in DMF (4ml) was added potassium fluoride (145mg, 2.5mmol, 2.5 equiv). After 10min stirring at room temperature, 2.6-dichlorobenzoic acid (229mg, 1.2mmol, 1.2 equiv) was added. After 3h 5 reaction at room temperature, ethyl acetate (30ml) was added. The solution was washed with a saturated solution of sodium bicarbonate (30ml), brine, dried over MgSO4 and concentrated in vacuo to afford 590mg (90%) of a pale yellow foam: $[\alpha]_n^{22}$ -85° (c 0.20. 10 CH₂Cl₂); IR (KBr) 3400, 2956, 1737, 1675, 1528, 1434, 1414, 1368, 1344, 1272, 1197, 1152, 1061; ¹H NMR $(CDCl_3) \delta 7.36-7.33$ (3H, m), 7.04 (1H, d, J = 8.0), 5.46 (1H, d, J = 7.8), 5.19-5.16 (1H, m), 5.08 (2H, AB),4.97 - 4.55 (1H, m), 4.69-4.55 (2H, m), 3.68 (3H, s), 15 3.30-3.10 (1H, m), 3.01-2.50 (4H, m), 2.40-2.33 (1H, m), 2.15-1.60 (5H, m), 1.44 (9H, s). Anal. Calcd for C28H34Cl2N4O10: C, 51.15; H, 5.21; N, 8.52. Found: C, 51.35; H, 5.32; N, 8.56.

[3S(1S,9S)] 5-(2,6-Dichlorobenzoyloxy)-3-[6,10-dioxo-9-20 (methoxycarbonyl) amino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxopentanoic acid (284), was synthesized from 508a via method used to prepare 505 from 504 which afforded 330mg (658) of a white solid: mp. 115°C (decomp.); 25 (α]_D²⁰ -107° (c 0.2, CH₂Cl₂); IR (KBr) 3340, 2954, 1736, 1664, 1530, 1434, 1272, 1198, 1148, 1060; ¹H NMR (D₆-DMSO) δ 8.91 (1H, d, J = 7.2H), 7.67-7.63 (3H, m), 7.54 (1H, d, J = 8.0), 5.24 (2H, s), 5.20-5.15 (1H, m), 4.79-4.70 (1H, m), 4.46-4.37 (2H, m), 3.58 (3H, s), 333-3,20 (1H, m), 2.94-2.55 (4H, m), 2.30-1.60 (6H.

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m). Anal. Calcd for $C_{24}H_{26}C_{12}N_4O_{10} \cdot H_2O$: C, 46.54; H, 4.56; N, 9.05. Found: C, 46.36; H, 4.14; N, 8.88.

[3S(1S,9S)] t-Butyl 5-(2,6-dimethylbenzoyloxy)-3-[6,10-dioxo-9-(methoxycarbonyl)amino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxopentanoate (508b), was synthesized by a similar method as compound 508a to afford a pale yellow foam (460mg, 82%): [α]_D²² -115° (c 0.20, CH₂Cl₂); IR (KBr) 3413, 2960, 1729, 1675, 1528, 1514, 1461, 1421, 1368, 1265, 1116, 1096;

1421, 1368, 1265, 1116, 1096;

14421, 1368, 1265, 1116, 1096;

15 NMR (CDCl₃) δ 7.27-7.03 (4H, m), 5.04 (2H, AB), 4.93-4.86 (1H, m), 4.80-4.56 (2H, m), 3.77 (3H, s), 3.32-3.15 (1H, m), 3.00-2.56 (4H, m), 2.37

(6H, s), 2.19-1.77 (5H, m), 1.45 (9H, s), 2.41-2.25 15 (1H, m). MS (ES⁺) 617.

30 559.

[3S(1S,9S)] 5-(2,6-Dimethylbenzoyloxy)3-[6,10-dioxo-9-(methoxycarbonyl)amino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxopentanoic acid (285), was synthesized by a similar method as compound 284 to afford a white solid (303mg, 78%): mp. 110°C (decomp.); [α]_D²⁰ -128° (c 0.10, CH₂Cl₂); IR (KBr) 3339, 2958, 1731, 1666, 1529, 1420, 1266, 1248, 1115, 1070; ¹H NMR (D₆-DMSO) δ 8.90 (1H, d, J = 7.4), 7.54 (1H, d, J = 7.9), 7.36-7.28 (1H, π., 2.717-7.14 (2H, m), 5.19-5.15 (3H, m), 4.84-4.74 (1H, m), 4.45-4.37 (2H, m), 3.59 (3H, s), 3.45-3.25 (1H, m), 2.95-2.64 (4H, m), 2.35 (6H, s), 2.30-1.60 (6H, m). Anal. Calcd for C₂₆H₃₂N₄O₁₀-H₂O: C, 53.98; H, 5.92; N, 9.68. Found: C, 53.50; H, 5.52; N, 9.49. MS (ES⁵)

509a-d

510a, 280, 283, 510d

compound	R
509a 510a	s—s
509b 280	s—N N N N N N N N N N N N N N N N N N N
509c 283	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
509d 510d	2 2 c

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[3S(1S,9S)] 3-(9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-5-(2-mercaptothiazole)-4-oxopentanoic acid (510a). A

dichloromethane (50ml) was treated with 30% hydrobromic

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acid in acetic acid (1.84ml, 9.2mmol, 2.2equiv) at 0°C, under nitrogen. After 10min stirring at 0°C the reaction was complete and a white solid crystallised in the medium. The solid was filtered and washed with 5 ethylacetate and diethylether to afford 2.20g (100%) of [3S(1S, 9S)] 5-bromo-3-(9-benzovlamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahvdro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4oxopentanoic acid which was used without further 10 purification: ¹H NMR (D₆-DMSO) δ 8.87 (1H, d, J = 7.3), 8.63 (1H, d, J = 7.6), 7.91-7.87 (2H, m), 7.60-7.44(3H, m), 6.92 (1H, bs), 5.14-5.09 (1H, m), 4.92-4.65 (2H, m), 4.43 (2H, AB), 4.41-4.35 (1H, m), 3.33-3.22 (1H, m), 2.98-2.90 (1H, m), 2.89-2.57 (2H, m), 2.35-15 2.15 (3H, m), 1.99-1.91 (2H, m), 1.75-1.60 (2H, m), A solution of the bromoketone (535mg, 1mmol) in dry DMF (10ml) was treated with potassium fluoride (150mg, 2.5mmol, 2.5 equiv), under nitrogen. After 5min stirring at room temperature, 2-mercaptothiazole 20 (140mg, 1.2mmol, 1.2equiv) was added. After overnight reaction ethylacetate (150ml) was added and the organic solution was washed with brine, dried over magnesium sulphate and reduced in vacuo. The residue was crystallised in diethyl ether, filtered and purified on 25 silica gel using a gradient of MeOH (0% to 5%) in dichloromethane. Evaporation afforded 344mg (60%) of a white solid: mp. 90-95°C (decomp.); $[\alpha]_{p}^{20}$ -82° (c 0.2, CH₂Cl₂); IR (KBr) 3328, 2941, 1745, 1659, 1535, 1422, 1276, 1255, 1223, 1072; ¹H NMR (D₆-DMSO) & 8.92 (1H, d. 30 J = 7.6), 8.68 (1H, d, J = 7.6), 7.98-7.90 (2H, m). 7.75-7.67 (1H, m), 7.64-7.50 (4H, m), 5.22-5.18 (1H. m), 4.95-4.74 (2H, m), 4.58-4.38 (3H, m), 3.52-3.19 (1H, m), 3.05-2.65 (4H, m), 2.40-1.50 (6H, m). Anal.

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Calcd for $C_{25}H_{27}N_5O_4S_2 \cdot H_2O$: C, 50.75; H, 4.94 N, 11.84. Found: C, 51.34; H, 4.70; N, 11.58. MS (ES⁺) 572.

[3S(1S,9S)] t-Butyl 3-(9-benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

- 5 pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-(1-phenyl-1H-tetrazole-5-thio) pentanoate (509b). 507a (100mg, 0.17mmol) in dry dimethylformamide (1.5ml) was treated with 1-phenyl-1H-tetrazole-5-thiol (33mg, 0.187mmol) and potassium fluoride (15mg, 0.34mmol).
- The mixture was stirred at room temperature for 2h, diluted with ethyl acetate, washed with aqueous sodium bicarbonate (x2), brine, dried (MgSO₄) and evaporated. The product was purified by flash chromatography on silica gel eluting with ethyl acetate to give 103mg
- 15 (88%) as a colourless foam: $\left[\alpha\right]_{D}^{23}$ -92.2° (c 0.1, CH₂Cl₂); IR (KBr) 3334, 1726, 1660, 1528, 1501, 1417, 1394, 1368, 1279, 1253, 1155; $^{1}_{1}$ H NMR (CDCl₃) 3 7.82 (2H, m), 7.60-7.40 (8H, m), 7.39 (1H, d, J = 8.1), 7.05 (1H, d, J = 7.3), 5.26 (1H, m), 5.15 (1H, m), 4.99 (1H, m),
- 20 4.60 (2H, m), 4.30 (1H, d, J = 17.2H), 3.32 (1H, m), 3.10-2.75 (4H, m), 2.40 (1H, m), 2.24 (1H, m), 1.90 (3H, m), 1.75 (1H, m), 1.44 (9H, s). MS (ES⁺) 691.47 (M⁺ + 1).

[3S(1S,9S)] 3-(9-Benzoylamino-6,10-dioxo-

25 1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxo-5(1-phenyl-1H-tetrazole-5-thio) pentanoic acid (280), was synthesized via method used to prepare 505 from 504. 509b (98mg, 0.142mmol) in dichloromethane (1ml) 30 was cooled to 0° and trifluoroacetic acid (1ml) was

added. The mixture was stirred at 0° for 15min and at

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room temperature for 30min before evaporation under reduced pressure. The residue was triturated with dry toluene and evaporated. Chromatography on silica gel eluting with 10% methanol in dichloromethane gave a colourless glass which was crystallised from dichloromethane/diethyl ether to give 62mg (69%) of colourless solid: mp. 145°C (decomp.); [α]_D²² -80.9° (c 0.1, CH₂Cl₂); IR (KBr) 3400, 1727, 1658, 1530, 1501, 1460, 1445, 1416, 1280, 1254; ¹H NMR (CDCl₃) δ 8.00 (1H, 10 m), 7.79 (2H, d, J = 6.7), 7.58-7.30 (9H, m), 5.25 (2H, m), 4.94 (1H, m), 4.53 (2H, m), 4.35 (1H, m), 3.35 (1H, m), 3.01 (3H, m), 2.73 (1H, m), 2.38 (1H, m), 1.98 (4H, m), 1.64 (1H, m). Anal. Calcd for C₂₉H₃₀N₈O₇S·0.2TFR: C, 53.71; H, 4.63 N, 17.04. Found: C, 53.97; H, 4.92; N, 15 16.77. MS (ES⁺) 633.55 (M⁺ - 1).

dioxo-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5(3-pyridyloxy)pentancate (509c), was prepared by a

20 similar method as compound 509b to afford a colourless
glass (34%): [α]_D²² -77.1° (c 0.25, CH₂Cl₂); IR (film)
3311, 1724, 1658, 1603, 1578, 1536, 1488, 1458, 1426,
1368, 1340, 1279, 1256, 1231, 1155, 707; ¹H NMR (CDCl₃)
δ 8.29 (2H, m), 7.84 (2H, m), 7.48 (4H, m), 7.22 (3H,
25 m), 5.20 (2H, m), 4.90 (2H, m), 4.58 (1H, m), 3.29 (1H,
m), 3.20-2.70 (4H, m), 2.38 (2H, m), 1.96 (4H, m), 1.68

[3S(1S,9S)] t-Butyl 3-[9-benzoylamino-6,10-

[3S(1S,9S)] 3-[9-Benzoylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-30 pvridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-

(1H, m), 1.42 (9H, s). MS (ES⁺) 608.54 (M + 1).

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(3-pyridyloxy)pentanoic acid (283), was prepared by a similar method as compound 280 to afford a colourless foam (100%): mp. ~125°C; [α]_D¹⁹ -84.1° (c 0.1, 20% MeOH/CH₂Cl₂); IR (KBr) 3401, 1736, 1663, 1538, 1489, 5 1459, 1425, 1281, 1258, 1200, 1134; ¹H NMR (CD₃OD/CDCl₃) δ 8.38 (2H, m), 7.84-7.40 (8H, m), 5.16 (4H, m), 4.80 (1H, m), 4.56 (1H, m), 3.50 (1H, m), 3.12 (2H, m), 2.82 (2H, m), 2.37 (1H, m), 2.10-1.65 (5H, m). Anal. Calcd for C₂₇H₂₉N₅O₈*0.4H₂O: C, 51.77; H, 4.61; N, 10.41. 10 Found: C, 52.19; H, 4.93; N, 9.99.

[3S(1S,9S)] t-Butyl 3-[6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-9-(phenycarbonylamino)-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-{2-[4(3H)-pyrimidone])pentanoate (509d), was

- 15 synthesized by a similar method as compound 509b to afford a colourless solid (49.6mg, 82%): ¹H NMR (CDCl₃)
 8.02 (1H, s), 7.95-7.86 (1H, m), 7.84-7.76 (2H, m), 7.62-7.35 (4H, m), 7.22-7.07 (1H, m), 6.43 (1H, d), 5.26-5.08 (2H, m), 5.03-4.72 (3H, m), 4.66-4.50 (1H, m), 3.43-3.19 (1H, m), 3.15-2.97 (1H, m), 2.86-2.72 (3H, m), 2.48-2.31 (1H, m), 2.18-1.60 (6H, m), 1.43 (9H, s).
 - [3S(1S,9S)] 3-[6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-9-(phenycarbonylamino)-6H-

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compound	R
504f 505f	CI N
504g 280b	N-N S-N
504h 283b	

15 [3s(1s,9s)] 5-(3-Chloro-2-oxy-4Hpyrido[1,2-a]pyrimidin-4-one)-3-[6,10-dioxo-9-

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(methylsulphonyl) amino-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4oxopentanoic acid (505f), was prepared by a similar
method as compound 508a using 507b and 3-chloro-2bydroxy-4H-pyrido[1,2-a]pyrimidin-4-one and directly
followed by the hydrolysis of 504f with trifluoroacetic
to afford a tan powder (65mg, 30%): [a]_D²⁰ -128° (c
0.10, MeOH); IR (KBr) 3414, 2928, 1667, 1527, 2459,
1407, 1328, 1274, 1153, 1134, ¹H NMR (MeOD) δ 9.35 (1H,
d, J = 6.6H), 8.34 (1H, t, J = 7.2H), 7.99-7.95 (1H,
m), 7.76-7.69 (1H, m), 5.85-5.45 (3H, m), 5.30-5.21
(1H, m), 4.93-4.66 (2H, m), 3.81-3.65 (1H, m), 3.66
(3H, m), 3.45-2.52 (4H, m), 2.52-1.71 (6H, m), D.J.

15 [3S(1S,9S)] t-Butyl 3-(6,10-dioxo-9-methanesulphonamido-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxo-5(1-phenyl-1H-tetrazole-5-thio)pentanoate (504g), was prepared by a similar method as compound 509b, (83%) as 20 a colourless foam: [α]_D²³ -112.7° (c 0.2, CH₂Cl₂); IR (KBr) 3312, 1726, 1668, 1501, 1413, 1395, 1369, 1328, 1276, 1254, 1155; ¹H NMR (CDCl₃) δ7.59 (5H, m), 7.48 (1H, d, J = 8.0), 5.68 (1H, d, J = 9.0), 5.37 (1H, m), 4.95 (1H, m), 4.62-4.31 (4H, m), 3.36 (1H, m), 2.98 (3H, s), 2.88 (4H, m), 2.66 (1H, m), 2.42 (2H, m., 1.98 (1H, m), 1.75 (1H, m), 1.43 (9H,s).

Hlasta et al., J. Med. Chem. 1995, 38, 4687-4692.

[3S(1S,9S)] 3-(6,10-Dioxo-9-methanesulphonamido-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxo-30 5(1-phenyl-1H-tetrazole-5-thio)pentanoic acid (280b),

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was prepared by a similar method as compound 280, (100%) as a colourless foam: mp. 120-5°C; $\left[\alpha\right]_D^{25}$ - 112.4° (c 0.1, CH₂Ci₂); IR (KBr) 3328, 1730, 1664, 1529, 1501, 1410, 1328, 1277, 1219, 1153, 1134, 991; 1_H 5 NMR (CDCl₃) δ 8.07 (1H, d, J = 7.8), 7.58 (5H, s), 6.41 (1H, d, J = 9.5), 5.32 (1H, m), 5.04 (1H, m), 4.70 (1H, d, J = 17.5), 4.60 (3H, m), 3.50-2.9 (3H, m), 2.98 (3H, s), 2.45 (2H, m), 2.06 (4H, m), 1.68 (1H, m).

[3S(1S,9S)] t-Butyl 3-(6,10-dioxo-9-

- 10 methanesulphonamido-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxo-5(3-pyridyloxy)pentanoate (504h), was prepared by a similar method as compound 509b (24%) as a colourless foam: [\alpha]r^{23} -101.0° (c 0.2, CH₂Cl₂); IR (KBr) 3330,
- 15 1727, 1669, 1425, 1396, 1369, 1328, 1276, 1256, 1231, 1155, 1137, 991; ¹H NMR (CDCl₃) &8.28 (2H, br d, J = 9.4), 7.71 (1H, d, J = 7.9), 7.22 (2H, s), 6.03 (1H, d, J = 9.4), 5.36 (1H, m), 4.95 (2H, m), 4.52 (2H, m), 3.29 (1H, m), 3.07 (3H, s), 3.23-2.75 (3H, m), 2.66-20 2.35 (2H, m), 2.30-1.60 (5H, m), 1.42 (9H, s).
 - [3S(1S,9S)] 3-(6,10-Dioxo-9-methanesulphonamido-1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5(3-pyridyloxy)pentanoic acid (283b), was prepared by a

25 similar method as compound 280, (100°) as a colourless foam: mp. 120-5°C; [α]_D²⁵-85.2° (c 0.1, 10% CH₃OH/CH₂Cl₂); IR (KBr) 3337, 1738, 1667, 1560, 1457, 1424, 1326, 1317, 1278, 1258, 1200, 1189, 1150, 1133, 991; ¹H NMR (CDCl₃/CD₃OD) δ 8.35 (2H, m), 7.54 (2H, m,

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5.32 (2H, m), 4.83 (2H, m), 4.45 (2H, m), 3.43-2.77 (4H, m), 2.97 (3H, s), 2.42 (2H, m), 2.05-1.72 (5H, m).

508c 511c SNN SOBE 280c SNN SOBE 283c

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[3S(1S,9S)] t-Butyl 3-[6,10-dioxo-9-(methoxycarbonyl)amino-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-5-(2-mercaptopyrimidine)-4-oxo-pentanoate (508c), was
15 prepared by a similar method as compound 509b to afford 544mg (97%) of a pale yellow foam: [\alpha_D^{20} -86° (c 0.19,

CH-Cl₂); IR (KBr) 3426, 2947, 1725, 1669, 1551, 1418,

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1383, 1253, 1155, 1064; 1 H NMR (CDC1₃) 8 8.49 (2H, d, J = 4.8), 7.13 (1H, d, J = 7.9), 7.03-6.98 (1H, m), 5.47 (1H, d, J = 7.9), 5.23-5.19 (1H, m), 5.09-5.01 (1H, m), 4.84-4.51 (2H, m), 4.04 (2H, AB), 3.69 (3H, s), 3.38-5 (3H, m), 3.06-2.64 (4H, m), 2.40-1.76 (6H, m), 1.43 (9H, s). Anal. Calcd for $C_{25}H_{34}N_{6}O_{8}S$: C, 51.89; H, 5.92; N, 14.52. Found: C, 51.49; H, 6.04; N, 13.87. MS (ES⁵) 579.

[3S(1S,9S)] 3-[6,10-Dioxo-9-(methoxycarbonyl)-amino10 1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a](1,2]diazepine-1-carboxamido]-5-(2mercaptopyrimidine)-4-oxopentanoic acid (511c), was
prepared by a similar method as compound 280 to afford
370mg (79%) of a white powder: mp. 105°C (dec); [α]_D²²
15 -94° (c 0.20, CH₂Cl₂); IR (KBr) 3316, 3057, 2957, 1724,
1664, 1252, 1416, 1384, 1254, 1189, 1063;

1H NMR (D₆DMSO) δ 8.85 (1H, d, J = 7.8), 8.62 (2H, d, J = 4.7),
7.53 (1H, d, J = 8.0), 7.28-7.23 (1H, m), 5.21-5.17
(1H, m), 4.87-4.79 (1H, m), 4.47-4.35 (2H, m), 4.23
20 (2H, AB), 3.58 (3H, s), 3.30-3.21 (1H, m), 2.95-2.50
(4H, m), 2.35-1.60 (6H, m). Anal. Calcd for
C₂₁H₂₆N₆O₈S·H₂O: C, 46.66; H, 5.22; N, 15.55. Found: C,
46.66; H, 5.13; N, 15.07. MS (ES[†]) 523. (ES[†]) 521.

[3S(1S,9S)] t-Butyl 3-[6,10-dioxo-9-

25 (methoxycarbonylamino)-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5[5-(1-phenyltetrazolyl)-thio]pentanoate (508d), was
synthesized by a similar method as compound 509b to
afford a colourless solid (269mg, 87*): mp. 80-110°C;
30 [a]n²³-108° (c 0.60 CH₂Cl₂); IR (KBr) 3315, 2977, 1727,

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1688, 1527, 1501, 1458, 1418, 1368, 1279, 1250, 1155,
1064; ^{1}H NMR (CDCl₃) δ 7.70 (1H, d), 7.63-7.53 (5H, m),
5.84 (1H, d), 5.34-5.27 (1H, m), 5.05-4.92 (1H, m),
4.78-4.54 (3H, m), 4.38 (1H, d), 3.66 (3H, s), 3.37-5
3.19 (1H, m), 3.07-2.94 (1H, m), 2.91-2.82 (2H, m),
2.71-2.56 (1H, m), 2.40-2.30 (1H, m), 2.19-2.13 (1H, m), 2.08-1.68 (4H, m), 1.42 (9H, s). MS (ES $^{+}$) 667 (318), 645 (M $^{+}$ + 1, 100), 589 (62).

[3S(1S,9S)] 3-[6,10-Dioxo-9-(methoxycarbonylamino)-

10 1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-[5-(1-phenyltetrazolyl)-thio]pentanoic acid (280c), was synthesized by a similar method as compound 280 to afford a pale cream solid (203mg, 88%): mp. 105-130°C; 15 [α]_D²² -235° (c 0.11 MeOH); IR (KBr) 3342, 2951, 1727, 1667, 1529, 1501, 1459, 1416, 1276, 1252, 1225, 1192,

1062; ^{1}H NMR (D₆-DMSO) δ 8.89 (1H, d), 7.69 (5H, s), 7.50 (1H, d), 5.18-5.11 (1H, m), 4.79-4.69 (1H, m),

4.57 (2H, s), 4.42-4.32 (1H, m), 3.54 (3H, s), 2.92-20 2.63 (3H, m), 2.21-1.82 (5H, m), 1.65-1.57 (1H, m). MS

(ES⁺) 587 (M - 1, 100%).

[3S(1S,9S)] t-Butyl 3-[6,10-dioxo-9-

(methoxycarbonylamino) -1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-

- 25 (3-pyridinyloxy) pentanoate (508e), was synthesized by a similar method as compound 509b to afford a pale orange solid (199mg, 25%): mp. 80-120°C; [α]_D²³ -69° (c 0.51 CH₂Cl₂); IR (KBr) 3333, 2978, 1726, 1669, 1576, 1536, 1478, 1426, 1368, 1277, 1253, 1232, 1155, 1064;
- 30 1 H NMR (CDCl₃) δ 8.41-8.18 (2H, m), 7.81 (1H, d), 7.26-

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7.20 (2H, s), 5.91 (1H, d), 5.24-5.16 (1H, m), 5.07-4.86 (3H, m), 4.81-4.51 (2H, m), 3.67 (3H, s), 3.34-3.16 (1H, m), 3.10-2.81 (3H, m), 2.72-2.54 (1H, m), 2.41-2.31 (1H, m), 2.07-1.62 (5H, m), 1.47 (9H s). MS 5 (ES⁺) 562 (M⁺ + 1, 100%), 506 (38).

[3S(1S,9S)] 3-[6,10-Dioxo-9-(methoxycarbonylamino)-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-(3-pyridinyloxy)pentanoic acid (283c), was synthesized 10 by a similar method as compound 280 to afford an offwhite powder (167mg, 98%): mp. 90-105°C; $[\alpha]_n^{22}$ -106° (c 0.11 MeOH); IR (KBr) 3325, 3070, 2956, 1669, 1544, 1423, 1256, 1199, 1133, 1062; 1 H NMR (D₆-DMSO) δ 8.95 (1H, d), 8.45-8.20 (2H, m), 7.53-7.45 (3H, m), 5.19-15 5.08 (3H, m), 4.70-4.62 (1H, m), 4.41-4.30 (2H, m), 3.53 (3H, s), 2.92-2.68 (3H, m), 2.22-2.06 (2H, m),

1.95-1.82 (2H, m), 1.63-1.53 (1H, m), MS (ES[†]) 506 (M[†] + 1, 100%).

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compound	R
512a 280d	SAN
512b 283d	

5

[3s(1s,9s)] t-Butyl 3-(9-acetamido-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2]diazepine-1-carboxamido]-4-oxo-5-(1-phenyl-1H-tetrazole-5-thio)pentanoate (512a), was

- 10 prepared by a similar method as compound 509b, to afford (83%) as a colourless foam: $(\alpha)_D^{23}$ -129.6° (c 0.1, CH₂Cl₂); IR (KBr) 3323, 1726, 1664, 1531, 1501, 1444, 1415, 1394, 1369, 1279, 1254, 1156; 1_H NMR (CDCl₃) δ 7.59 (5H, s), 7.37 (1H, d, J = 7.9), 6.38 (1H,
- 15 d, J = 7.4), 5.27 (1H, m), 4.98 (2H, m), 4.58 (2H, d + m), 4.28 (1H, d, J = 17.2), 3.26 (1H, m), 3.10-2.65 (4H, m), 2.31 (2H, m), 2.03 (3H, s), 2.10-1.72 (4H, m), 1.48 (9H, s).

[3S(1S,9S)] 3-(9-Acetamido-6,10-dioxo-1,2,3,4,7,8,9,10-

20 octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1carboxamido)-4-oxo-5-(1-phenyl-1H-tetrazole-5thio)pentanoic acid (280d), was prepared by a similar
method as compound 280, to afford (77%) as a colourless
foam: [αIn²² -93.3° (c 0.1, CHaClo); IR (KBr) 3316,

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1728, 1659, 1531, 1501, 1415, 1341, 1278, 1253, 1222,
1185; ^{1}H NMR (CDCl $_{3}$) δ 8.05 (1H, d, J = 7.9), 7.57 (5H, br s), 5.30 (1H, m), 5.01 (2H, m), 4.70-4.10 (4H, m),
3.40-2.85 (4H, m), 2.62 (1H, m), 2.33 (1H, m), 2.27-5 1.65 (5H, m), 2.01 (3H, s).

[3S(1S,9S)] t-Butyl 3-(9-acetamido-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxo-5-(3-pyridyloxy)pentanoate (512b), was prepared by a similar method as compound 509b, to afford (9%) as a colourless foam: IR (KBr) 3333, 1727, 1661, 1542, 1427, 1369, 1279, 1257, 1232, 1156; ¹H NMR (CDCl₃) & 8.30 (2H, m), 7.20 (3H, m), 6.45 (1H, d, J = 7.4), 5.17 (1H, m), 4.91 (3H, m), 4.55 (1H, m), 3.27 (1H, m), 3.14-2.70 (4H, m), 2.41 (1H, m), 2.04 (3H, s), 2.10-1.65 (6H, m), 1.44 (9H, s).

[3S(1S,9S)] 3-(9-Acetamido-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxo-5-(3-pyridyloxy)pentanoic acid

20 (283d), was prepared by a similar method as compound 280. (100%) as colourless foam: [\alpha]_0^22 -106.0° (c 0.2, 10% CH₃OH/CH₂Cl₂); IR (KBr) 3312, 1735, 1664, 1549, 1426, 1279, 1258, 1200, 1135; \frac{1}{14} NMR (CDCl₃) \delta 8.27 (2H, m), 7.46 (2H, m), 5.09 (1H, m), 4.79 (3H, m), 4.47 (1H, 25 m), 3.40 (1H, m), 3.30-2.70 (3H, m), 2.54 (1H, m), 2.35 (1H, m), 1.98 (3H, s), 2.05-1.65 (4H, m).

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245b 246b

[15,9R(2RS,3S)] 9-Benzoylamino-N-(2-benzyloxy-5-oxotetrahydrofuran-3-yl)-1,2,3,4,7,8,9,10-octahydro-10-oxo-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamide

- 5 (245b), was prepared from (1S,9R) 9-Benzoylamino-1,2,3,4,7,8,9,10-octahydro-10-oxo-6Hpyridazino[1,2-a][1,2]diazepine-1-carboxylic acid by the method described for 245 to afford 416mg (85%) of a colourless foam (~1:1 mixture of diastereoisomers): IR
- 10 (KBr) 3392, 3302, 2942, 1792, 1642, 1529, 1520, 1454,
 1119; ^{1}H NMR (CDCl $_{3}$) δ 7.79 (2H, m), 7.51-7.09 (10H, m),
 5.52 (0.5H, d, J = 5.3), 5.51 (0.5H, s), 5.36 (1H, m),
 4.84 (1H, m), 4.74-4.59 (1.5H, m), 4.51 (1H, m), 4.38 (0.5E, m), 3.22-2.83 (5H, m), 2.51 (1H, m), 2.25 (2H,
- 15 m), 2.01-1.46 (6H, m). Anal. Calcd for $C_{28}H_{32}N_4O_6 \cdot 0.75H_2O$: C, 62.97; H, 6.32; N, 10.49. Found: C, 63.10; H, 6.16; N, 10.21. MS (ES⁺) 521 (M + 1, 100%).

[3S(1S,9R)] 3-(9-Benzoylamino-1,2,3,4,7,8,9,10-

20 octahydro-10-oxo-6H-pyridazino[1,2-a][1,2]diazepine-1-carboxamido)-4-oxobutanoic acid (246b), was prepared from 245b by the method described for 246 to afford

104mg (33%) of a white powder: mp. 115-119°C; ${\left(\alpha\right)_{D}}^{24}$ - 19.8° (c 0.2 MeOH); IR (KBr) 3293, 2944, 1786, 1639, 1576, 1537, 1489, 1450, 1329, 1162, 1124; 1 H NMR (CD₃OD) δ 7.85 (2H, d, J = 7.0), 7.49 (3H, m), 5.49 (1H, 5 m), 4.55 (1H, m), 4.30 (2H, m), 3.40 (1H, m), 3.19-2.89 (3H, m), 2.63 (2H, m), 2.16-1.81 (5H, m), 1.60 (3H, m). Anal. Calcd for C₂₁H₂₆N₄O₆*H₂O: C, 56.24; H, 6.29; N, 12.49. Found: C, 56.54; H, 6.05; N, 12.29. MS (ES[†]) 429 (M - 1, 100%).

10 Compounds 513a-j were prepared as described below.

513a-f

compound	R
513a	,°~~
5 13a-1	~°~~
513a-2	
513b	хо <u>(</u>)
513b-1	·

15

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513b-2	,o-<
513c	`o-(\(\)()
513d	,,o,o,,
513e	,°~
513f	D 0 √
513f-1	"·°~
513f-2	<i>^</i> ~

5

513g

(2RS,3S) 3-(Allyloxycarbonyl)amino-2-(2-phenethyloxy)5 5-oxotetrahydrofuran (513a), was prepared by a similar method as compound 513d/e to afford a mixture of diastereoisomers (670mg, 50%) as an oil: IR (KBr) 3331, 2946, 1790, 1723, 1713, 1531, 1329, 1257, 1164, 1120, 1060, 977, 937, 701; ¹H NMR (CDCl₃) δ 7.36-7.18 (5H, m), 5.99-5.83 (1H, m), 5.41-5.34 (2H, m), 5.28-5.18 (2H,

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m), 4.59-4.56 (2H, m), 4.32-3.96 (2H, m), 3.85-3.73 (1H, m), 3.02-2.76 (3H, m), 2.49-2.34 (1H, m).

 $(2RS,3S) \ \ 3-(Allyloxycarbonyl) \ amino-2-cyclopentyloxy-5-oxotetrahydrofuran (513b), was prepared as 513d/e to afford 8g (518) of a mixture of diastereoisomers as a clear oil: <math>[\alpha]_D^{2C}-13^\circ$ (c 0.25, CH_2Cl_2); IR (KBr) 3325, 2959, 2875, 1790, 1723, 1535, 1420, 1328, 1257, 1120, 1049, 973, 937; 1H NMR (CDCl₃) δ 6.02-5.80 (1H, m), 5.53-5.46 (2H, m), 5.37-5.21 (2H, m), 4.58 (2H, d, J = 5.5), 4.50-4.46 (0.5H, m), 4.34-4.25 (1H, m), 4.19-4.12 (0.5H, m), 3.06-2.77 (1H, m), 2.53-2.35 (1H, m), 1.85-1.50 (8H, m). Anal. Calcd for $C_{13}H_{19}NO_5$: C, 57.98; H, 7.11; N, 5.20. Found: C, 56.62; H, 7.22; N, 4.95. MS

15 (2R,38) 3-Allyloxycarbonylamino-2-(indan-2-yloxy)-5-oxotetrahydrofuran (513c), was synthesized by a similar method as compound 513d/e to afford a single isomer (20%) as a pale yellow oil: [α]_D²⁴ -63.1° (c 0.2, CH₂Cl₂); IR (film) 3338, 2948, 1791, 1723, 1529, 1421,
 1330, 1253, 1122, 984, 929, 746; ¹H NMR (CDCl₃) 87.20 (4H, m), 5.87 (1H, m), 5.61 (1H, d, J = 5.4), 5.33-5.10 (2H, m), 4.70 (1H, m), 4.56 (3H, m), 3.33-3.19 (2H, m), 3.10-2.94 (2H, m), 2.81 (1H, dd, J = 8.3, 17.3), 2.43 (1H, dd, J = 10.5, 17.3).

(ES⁺) 270.

25 (2R,3S) 3-Allyloxycarbonylamino-2-benzyloxy-5-oxotetrahydro-furan (513d) and (2S,3S) 3-Allyloxycarbonylamino-2-benzyloxy-5-oxo-tetrahydrofuran (513d/e), were prepared [via method described by Chapman Biorg. & Med. Chem. Lett., 2, pp. 615-618

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(1992)]. Following work-up by extraction with ethylacetate and washing with NaHCO3, the product was dried (MgSO4), filtered and evaporated to yield an oil which contained product and benzyl alcohol. Hexane

- 5 (200ml) (200ml hexame for every 56g of AllocAsp(CO₂tBu)CH₂OH used) was added and the mixture stirred and cooled overnight. This afforded an oily solid. The liquors were decanted and retained for chromatography. The oily residue was dissolved in
- 10 ethyl acetate and evaporated to afford an oil which was crystallised from 10% ethyl acetate in hexane (~500ml). The solid was filtered to afford 513d (12.2q, 19%): mp. $108-110^{\circ}\text{C}$; [α] $_{0}^{24}$ +75.72° (c 0.25, CH₂Cl₂); IR (KBr)3361, 1778, 1720, 1517, 1262, 1236, 1222, 1135,
- 15 1121, 944, 930, 760; ¹H NMR (CDCl₃) 87.38 (5H, m), 5.90 (1H, m), 5.50 (1H, s), 5.37 (0.5H, m), 5.26 (2.5H, m), 4.87 (1H, ABq), 4.63 (3H, m), 4.31 (1H, m), 3.07 (1H, dd), 2.46 (1H, dd). Anal. Calcd for C₁₅H₁₇NO₅: C, 61.85; H, 5.88; N, 4.81. Found: C, 61.85; H, 5.89; N, 4.80.
- The liquors were combined and evaporated to yield an oil (~200g) containing benzyl alcohol. Hexane/ethyl acetate (9:1, 100ml) was added and the product purified by chromatography eluting with 10% ethyl acetate in hexane to remove the excess benzyl alcohol, and then dichloromethane/hexane (1:1 containing 10% ethyl acetate). This afforded 513e containing some 513d (20.5g, 32%): mp. 45-40°C; [\alpha]_0^C4 -71.26° (c 0.25, CH2Cl2); IR (KBr) 3332, 1804, 1691, 1536, 1279, 1252, 1125,976. H NMR (CDCl3) \delta 7.38 (5H, 30 m), 5.91 (1H, m), 5.54 (1H, d, J = 5.2), 5.38 (3H, m);

4.90 (1H, ABq); 4.60 (4H, m), 2.86 (1H, dd); 2.52 (1H,

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dd). Anal. Calcd for $C_{15}H_{17}NO_5 \cdot 0.1H_2O$ C, 61.47; H, 5.91; N, 4.78. Found: C, 61.42; H, 5.88; N, 4.81.

(2RS,3R) 3-(Allyloxycarbonylamino)-2-ethoxy-5oxotetrahydrofuran (513f), was synthesized by a similar

5 method as 513d/e to afford a colourless oil (152mg,
79%): IR (film) 3334, 2983, 2941, 1783, 1727, 1713,
1547, 1529, 1422, 1378, 1331, 1313, 1164, 1122, 1060,
938; ¹H NMR (CDCl₃) & 6.09-5.82 (2H, m), 5.50-5.18 (3H,
m), 4.64-4.54 (2H, m), 4.27-4.16 (1H, m), 3.95-3.78

10 (1H, m), 3.73-3.56 (1H, m), 3.05-2.77 (1H, m), 2.562.37 (1H, m), 1.35-1.17 (4H, m). Anal. Calcd for
Cl₁₀H₁₅NO₅: C, 52.40; H, 6.60; N, 6.11. Found: C, 52.16;
H, 6.62; N, 5.99. MS (ES[†]) 229 (M[†] + 1, 100%).

(3S,4RS) t-Butyl 3-(allyloxycarbonylamino)-4-hydroxy-5-

- 25 (2-phenoxybenzoyloxy)pentanoate (513g). 4Dimethylamino-pyridine (76.0mg, 622mmol) was added to a solution of 2-phenoxybenzoyl chloride (579mg, 2.49mmol) and 517 (600mg, 2.07mmol) in pyridine (10ml). The mixture was stirred at room temperature for 18h before 20 adding brine (25ml) and extracting with ethyl acetate (30ml, 20ml). The combined organic extracts were washed with 1M hydrochloric acid (3 x 25ml), saturated aqueous sodium hydrogen carbonate (2 x 25ml) and brine (25ml), dried (MqSO_A) and concentrated. The pale

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(3S,4R) t-Butyl (N-allyloxycarbonyl)-3-amino-4-hydroxy-5-(1-naphthoyloxy)pentanoate (513h), was prepared from (3S,4R) t-butyl (N-allyloxycarbonyl)-3-amino-4,5
10 dihydroxypentanoate by the method described for 513g to afford 562mg (85%) of a colourless oil: IR(film) 3418, 2980, 1722, 1711, 1512, 1368, 1278, 1245, 1198, 1157, 1139;

1 h NMR (CDCl₃) & 8.90 (1H, d, J = 8.6), 8.21 (1H, dd, J = 1.2, 7.3), 8.04 (1H, d, J = 8.2), 7.89 (1H, dd, 15 J = 1.5, 7.9), 7.67-7.46 (3H, m), 5.86 (1H, m), 5.49 (1H, d, J = 9.0), 5.35-5.18 (2H, m), 4.57-4.46 (4H, m), 4.19 (2H, m), 2.67 (2H, m), 1.40 (9H, s). Anal. Calcd for C₂₄H₂₉NO₇: C, 65.00; H, 6.59; N, 3.16. Found: C, 64.74; H, 6.56; N, 3.09. M.S. (ES⁺) 466 (M+Na, 100)),

20 444 (M+1, 39), 388 (44).

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486 (M $^+$ + 1, 33. Accurate mass calculated for $C_{2.6}H_{3.2}NO_{\theta}$ (M \dot{H}^+): 486.2128. Found: 486.2121.

(35,4RS) t-Butyl 3-(allyloxycarbonylamino)-4-hydroxy-5-(5-methyl-3-phenylisoxazoloyloxy)pentanoate (513j), was 5 synthesized by a similar method as compound 513g to afford a pale orange oil (905mg, 91%): IR (film) 3418, 3383, 2980, 1722, 1711, 1601, 1517, 1450, 1424, 1368, 1308, 1252, 1154, 1100, 994, 767, 6981 H NMR (CDCl₃) & 7.62-7.55 (2H, m), 7.51-7.42 (3H, m), 5.98-5.76 (1H, 10 m), 5.33-5.18 (2H, m), 4.53 (2H, d), 4.18 (2H, d), 3.91 (1H, m), 3.80 (1H, m), 2.76 (3H, s), 2.50 (2H, m), 14.3 (9H, s). Anal. Calcd for C₂₄H₃₀N₂O₈*0.5H₂O: C, 59.62; H, 6.46; N, 5.79. Found: C, 59.46; H, 6.24; N, 5.72. MS (ES⁺) 497 (100%), 475 (M⁺ + 1, 15), 419 (48).

15

(35,4R) t-Butyl 3-benzylamino-4,5-(dimethylmethylenedioxy)-pentanoate (514), was prepared by the method described in H. Matsunaga, et al. Tetrahedron Letters 24, pp. 3009-3012 (1983) as a pure diastereomer (60%) as an oil: $\left[\alpha\right]_{D}^{23}$ -36.9° (c 0.5, dichloromethane); IR (film) 2982, 2934, 1726, 1455, 1369, 1257, 1214, 1157, 1068; ^{1}H NMR (CDCl₃) δ 7.31 (5H, 5 m), 4.10 (1H, q, J = 6.0), 4.05-3.75 (4H, m), 3.10 (1H, q, J = 6.0), 2.40 (2H, m), 1.42 (9H, s), 1.40 (3H, s), 1.34 (3H, s).

(3s,4R) t-Butyl 3-(allyloxycarbonylamino)-4,5-(dimethylmethylenedioxy)pentanoate (516). 514 (3.02g,

- 9.00mmol) and 10% palladium on carbon (300mg) in ethanol (30ml) were stirred under hydrogen for 2h. The suspension was filtered through celite and a 0.45mm membrane and the filtrate concentrated to give a colourless oil 515 (2.106g, 95%) which was used without 15 purification. The oil (1.93g, 7.88mmol) was dissolved
- 15 purification. The oil (1.93g, 7.88mmol) was dissolved in water (10ml) and 1,4-dioxan and sodium hydrogen carbonate added (695mg, 8.27mmol). The mixture was cooled to 0°C and allyl chloroformate (1.04g, 919ml, 8.66mmol) added dropwise. After 3h the mixture was
- 20 extracted with ether (2 x 50ml). The combined ether extracts were washed with water (2 x 25ml) and brine (25ml), dried (MgSO $_4$) and concentrated to give a colourless oil. Flash column chromatography (10-35* ethylacetate in hexane) afforded a colourless solid
- 25 (2.69g, 95%): mp. 64-5°C; $(\alpha)_D^{23}$ -21° (c 1.00, CH2Cl2.); IR (KBr) 3329, 1735, 1702; ^{1}H NMR (CDCl2) δ 6.00-5.82 (1H, m), 5.36-5.14 (2H, m), 542 (1H, s), 4.56 (1H, d), 4.40-4.06 (2H, m), 4.03 (1H, m) 3.70 (1H, m), 2.52 (2H, m), 1.44 .12H, 2 x s), 1.33 (3H, s); Anal. Calcd for
- 30 C₁₆H₂₇NO₆: C, 58.34; H, 8.26; N, 4.25. Found : C, 58.12; H, 8.16; N, 4.19; MS (+FAB) 320 (M⁺+1, 41|1), 274 (70), 216 (100).

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(35,4R) t-Butyl 3-(allyloxycarbonylamino)4,5-dihydroxy pentanoate (517). A solution 516 (2.44g, 7.41mmol) in 80% aqueous acetic acid (25ml) was stirred at room temperature for 24h then concentrated and azeotroped

- 5 with toluene (2 x 25ml). The residue was treated with brine (25ml) and extracted with ethylacetate (2 x 25ml). The organic fractions were dried (MgSO₄) and concentrated to afford a colourless oil. Flash chromatography (20-80% ethyl acetate in
- 10 dichloromethane) gave a colourless solid (1.99g, 90%): mp. $74-5^{\circ}\text{C}$; [α] $_{D}^{25}-1.3^{\circ}$ (c 1.0, $\text{CH}_{2}\text{Cl}_{2}$); IR (KBr) 1723, 1691; ^{1}H NMR (CDCl $_{3}$) δ 6.02-5.78 (2H, m), 5.35-5.16 (2H, m), 4.55 (2H, d), 4.16-4.04 (2H, m), 2.76 (2H, s), 3.56 (2H, m), 2.56 (2H, m), 1.43 (9H, s); Anal. Calcd
- 15 for $C_{13}H_{23}NO_6$: C, 53.97; H, 8.01; N, 4.84. Found : C, 53.79; H, 7.88; N, 4.81; MS(+FAB) 290 (M^+ +1, 44%), 234 (100).

Example 30

Compounds 1105-1125 were prepared as follows.

20 Physical data for these compounds is listed in Table 24.

MS (M+Na) +	496.9	496.9
HPLC RT min (method) Purity	12.769 (1)	12.137 (1)
MW	473.49	473.45
MF	C22H27N5O7	C21H23N508
Structure		
Compound	1105	1106

Table 24

MS (M+Na)+	502.9	536.4
HPLC RT min (method) Purity	11.272 (1) 97%	13.699 (1)
ММ	479.47	512.48
χ H	C19H21N5OBS	C23H24N60B
Structure	T I O Z I O S I S I S I S I S I S I S I S I S I	
Compound	1107	1108

Jilo Structure MF MW (method) (M+Na)+ 1109				
Structure MF MW C22H23N5010 517.46 MP C22H25N509 503.47	MS (M+Na)+	541.2	527.9	526.7
Structure MF CZ2H23NS010 CZ2H23NS010 CZ2H2SNS09 CZ2H2SNS09	HPLC RT min (method) Purity	12.341 (1)	12.991 (1) 96%	10.951 (1) 99%
Structure Structure Structure Structure Structure Structure Structure	MM	517.46	503.47	503.47
Structure Structure Structure Structure Structure Structure	M	C22H23N5O10	C22H25N509	C22H25N509
1109 1110	Structure	0= -0 ZI 0 Z-Z 00	0= ZI 0= ZI 0= ZI	0 2-2 0
	Compound	1109	1110	1111

MS (M+Na)+	557.2	531.5
HPLC RT min (method) Purity	98%	16.317 (1)
ММ	533.50	507.93
MF	C23H27N5O10	C2ZH26CINSO7 507.93
Structure	O N N DOH	0 Z Z 0 0 Z I
Compound	1112	1113

		T
MS (M+Na)+	542.4	563.4
HPLC RT min (method) Purity	12.902 (1) 99%	12.529 (2) 97%
MM	517.50	540.36
ΜF	C23H27N509	C22H23C12N5O7 540.36
Structure		HO H HO N H
Compound	1114	1115

MS (M+Na)+	538.8	538.8
HPLC RT min (method) Purity	14.144 (1)	11.551 (2) 97%
MW	515.48	515.53
MF	C23H25N509	C24H29N5OB
Structure	O N N O H	0 Z Z Z 0 0 Z I 0 0 Z I 0 0 0 0 0 0 0 0
Compound	1116	1117

MS (M+Na) + (84.9		4 8 9. 9.	502.9
HPLC RT min (method)	Purity	13.974 (1)	11.079 (2)
MW		465.51	479.54
Σi		C21H31N5O7	C22H33N5O7
Structure		D Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	
Compound		1118	1119

MS (M+Na)+	547.3	527.9
HPLC RT min (method)	16.796 (1)	11.131 (1)
ММ	522.91	503.47
MF	C21H23ClN6O8	C22H25N509
Structure	TI OZZI	
Compound	1120	1121

+	S	
MS (M+Na)+	525.5	574
HPLC RT min (method) Purity	10.892 (2) 98%	15.85
MM	501.54	552.50
MF	C24H31N5O7	C26H24N4010
Structure		Z-Z-O
Compound	1122	1123

				HPLC RT min	SW.
	Structure	Æ	MM	(method)	(M+Na)+
€-0 £		C24H29N5O11	563.53	13.336 (1)	587
5 5 5 7		C21H23C12N5O8 544.35	544.35	8 0 9 9 9 5	0 0

Step A. Synthesis of 401. TentaGel S% $\rm NH_2$ resin (0.25 mmol/g, 5.25 g) was placed in a sintered glass shaker vessel and washed with dimethylacetamide (3 X 15 mL). Compound 400 (1.36 g, 2.3 mmol) was

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dissolved in DMA (10 mL) and O-benzotriazole-N,N,N,N'tetramethyluronium hexafluorophosphate (HBTU; 0.88 g,
2.3 mmol), and DIEA (0.8 mL, 4.6 mmol) were added. The
solution was transferred to the resin and a further 5
mL DMA added. The reaction mixture was agitated for
1.5 h at room temperature using a wrist arm shaker.
The resin was filtered and washed with
dimethylacetamide (4 X 15 mL).

Step B. Synthesis of 1102. Resin 401 was

- 10 deprotected with 20% (v/v) piperidine/dimethylacetamide (15 mL) for 10 min (shaking) and then for 10 min with fresh piperidine reagent (15 ml). The resin was then washed with dimethylacetamide (6 X 15 ml), followed by N-methypyrrolidone (2 X 25 mL).
- Compound 1101 (0.979 g, 2.11 mmol) was dissolved in dimethylacetamide (8 mL). HBTU (0.81 g, 2.1 mmol) and DIEA (0.75 mL, 4.3 mmol) were added and the solution added to the resin, followed by dimethylacetamide (4 mL). The reaction mixture was agitated for 2 h at room temperature using a wrist arm shaker. The resin work-up was performed as described for 401 to yield 1102.

Step C. Synthesis of 1103. This compound was prepared from resin 1102 (0.040 mmol) using an Advanced ChemTech 396 Multiple Peptide synthesizer. The automated cycles consisted of a resin wash with dimethylformamide (2 X 1 mL), deprotection with 25% (V/V) piperidine in dimethylformamide (1 mL) for 3 min followed by fresh reagent (1 mL) for 10 min to yield 30 resin 1103. The resin was washed with

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dimethylformamide (3 X 1 mL) and N-methypyrrolidone (3 X 1 mL).

Resin 1103 was acylated with a solution of 0.4M carboxylic acid and 0.4M HOBT in N-5 methypyrrolidone (0.5 mL), a solution of 0.4M HBTU in N-methylpyrrolidone (0.5 mL) and a solution of 1.6M DIEA in N-methypyrrolidone (0.25 mL) and the reaction was shaken for 2 hr at room temperature. The acylation step was repeated. Finally, the resin was washed with 10 N-methylpyrrolidone (1 X 1 mL), dimethylformamide (4 X 1 mL), dichloromethane (5 X 1 mL) and dried in vacuo. The aldehyde was cleaved from the resin and globally deprotected by treatment with 95% TFA/ 5% H2O (v/v, 1.5 mL) for 30 min at room temperature. After washing the 15 resin with cleavage reagent (1 mL), the combined filtrates were added to cold 1:1 ether:hexane (10 mL) and the resulting precipitate was isolated by centrifugation and decantation. The resulting pellet was dissolved in 10% acetonitrile/90% HoO/0.1% TFA (5 20 mL) and lyophilized to obtain crude 1105-1125 as a white powder. The compound was purified by semipreparative RP-HPLC with a Rainin Microsorb C18 column (5 μ, 21.4 X 250 mm) eluting with a linear acetonitrile gradient (8% - 48%) containing 0.1% TFA (v/v) over 30 25 min at 12 mL/min. Fractions containing the desired product were pooled and lyophilized to provide 1105-

1125 (10.8 mg, 63%).

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Analytical HPLC methods:

- (1) Waters DeltaPak C18, 300Å (5 μ , 3.9 X 150 mm). Linear acetonitrile gradient (0% 25%) containing 0.1% TFA (v/v) over 14 min at 1 mL/min.
- 5 (2) Waters DeltaPak C10, 300Å (5µ, 3.9 X 150 mm). Linear acetonitrile gradient (5% - 45%) containing 0.1% TFA (v/v) over 14 min at 1 mL/min.

Benzyl 3-(N'-t-butyloxycarbonylhydrazino)propionate
(259b), was synthesized via method used to prepare 259

10 from 258 to afford a waxy solid (87g, 51): mp 54-55°C;

IR (film) 3324, 2978, 1732, 1713, 1455, 1367, 1277,

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1254, 1171; 1 H NMR (CDCl₃) δ 7.35 (5H, m), 6.15 (1H, bs), 5.13 (2H, s), 3.15 (2H, t, J = 6.5), 2.54 (2H, t, J = 6.5), 1.45 (9H, s). Anal. Calcd for $C_{15}H_{22}N_{2}O_{3}$: C, 61.21; H, 7.53; N, 9.52. Found: C, 61.29; H, 7.51; N, 5 9.51. MS (ES⁺) 295 (M⁺ + 1).

(3s) 1-Benzyl 3-t-butyl 2-(N-2-benzyloxycarbonylethyl-NI-2-butoxycarbonylhydrazino) carbonyl hexahydropyridazine dicarboxylate (260b), was synthesized via method used to prepare 260 from 259 to afford a qum (81g) which was used in the next step without purification. Analytical data for a pure sample: IR (film) 3318, 2976, 1733, 1451, 1412, 1393, 1366, 1256, 1161; ¹H NMR (CDCl₃) & 7.34 (10H, m), 6.68 (0.5H, bs), 5.11 (4H, m), 4.63 (0.5H, bs), 4.14 (1H, 15 m), 3.53 (2H, m), 3.08 (1H, m), 2.63 (2H, m), 2.10-1.60 (4H, m), 1.60-1.35 (19H, m + 2 x s).

(3s) t-Butyl 2-(N'-t-butoxycarbonyl-N-2-carboxyethylhydrazino)-carbonylhexahydropyridazine 3-carboxylate (261b), was synthesized via method used to prepare 261 from 260 to give a gum which was purified by flash chromatography (1:1 ethyl acetate/dichloromethane) to give the title compound 261b (36.0g, 79.4% over 2 stages): TR (film) 3267, 2979, 2937, 1728, 1668, 1394, 1369, 1245, 1159; H NMR 25 (CDCl₃) & 7.6 (1H, bs), 6.8 (1H, vbs), 4.47 (1H, bs), 3.73 (2H, bs), 2.98 (1H, bs), 2.66 (3H, m), 2.04 (1H, bs), 1.84 (1H, m), 1.6-1.2 (21H, m + s).

(4S) t-Butyl 7-t-butoxycarbonylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

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pyridazino[1,2-a][1,2,4]triazepine-4-carboxylate (262b), was synthesized via method used to prepare 262 from 261 to give the title compound 262b, (18.6g, 54%) as an oil: $[\alpha]_D^{20}$ +47.7° (c 0.236, CH₂Cl₂); IR (film) 5 3291, 2978, 1738, 1727, 1690, 1678, 1439, 1243, 1164; ¹H NNR (CDCl₃) &6.59 (1H, s), 5.06 (1H, m), 4.47 (1H, m), 3.85 (3H, m), 2.82 (1H, m), 2.37 (1H, m), 2.22 (1H, m), 1.92 (1H, m), 1.63 (2H, m), 1.48 and 1.46 (18H, 2 x s). MS ($\mathbb{E}\mathbf{S}^+$) 399 (M⁺ + 1).

- 10 (4S) t-Butyl 7-amino-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4carboxylate (518). Compound 262b (2.43g, 6.1mmol) was dissolved in 1M hydrogen chloride in ethyl acetate (30ml) and stirred at room temperature for 20h. Solid 15 sodium bicarbonate (4q, 46.5mmol) and water 20ml were added and the mixture stirred for 5min before separating and extracting the aqueous portion with ethyl acetate. The combined organic solution was washed with water, saturated salt, dried (MgSO4) and 20 concentrated. Purification by flash chromatography (50% ethyl acetate in dichloromethane - 100% ethyl acetate) gave the pure product 518 (1.08g, 59%) as an unstable oil: $[\alpha]_D^{20}$ +82° (c 0.55, CH₂Cl₂); IR (film) 3331, 2977, 1731, 1680, 1664, 1439, 1420, 1315, 1158; 25 ¹H NMR (CDCl₃) δ5.08 (1H, m), 4.48 (1H, m), 3.80 (2H,
- Abg; 3.70 (2H, bs, exch with D₂O), 3.53 (1H, m), 2.75 (1H, m), 2.30 (2H, m), 1.88 (1H, m), 1.71 (2H, m), 1.47 (9H, s).

carboxylate (520). 519 (9.4g, 35.6mmol) was suspended in methanol (230ml) and cooled to 0°C in an ice bath. Thionyl chloride (3ml, 4.89g, 41.1mmol) was added 5 dropwise over 30min and the mixture stirred at ambient temperature for 48h. The solvent was removed in vacuo at 30°C and the oily residue dissolved in ethyl acetate (500ml). The organic solution was washed with saturated sodium bicarbonate, water and brine, dried 10 (MgSO₄) and concentrated to give 520 (7.84g, 79°) as an oil: [\alpha]_2^2 -25.9° (c 0.615, CH₂Cl₂); IR (film) 2953, 1739, 1703, 1694, 1440, 1403, 1357, 1261, 1241, 1174;

¹H NMR (CDCl₃) \(\delta 7.36 \) (5H, s), 5.18 (2H, s), 4.00 (1H,

(3S) Methyl 1-benzyloxycarbonyl-hexahydropyridazine-3-

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bd), 3.73 (3H, s), 3.55 (1H, dd), 3.12 (1H, t), 2.06 (1H, m), 1.73 (3H, m). Anal. Calcd for $C_{14}H_{17}N_{2}O_{4} \cdot 0.25H_{2}O$: C, 59.46; H, 6.59; N, 9.91. Found: C, 59.44; H, 6.46; N, 10.09.

- 5 (3S) 1-Benzyl 3-methyl 2-(N-2-benzyloxycarbonylethyl-NI-t-butoxycarbonylhydrazino)carbonyl hexahydropyridazine dicarboxylate (521). Using a similar method to that described for 260 above, 521 was prepared, 96% as a crude oil: [α]_D²² -22.16° (c 0.25, 10 CH₂Cl₂); IR (film) 3316, 2976, 2953, 1738, 1726, 1714, 1690, 1367, 1260, 1167; ¹H NMR (CDCl₃) δ7.25 (10H, m), 6.82 (1H, bs), 5.10 (4H, m), 4.80 (1H, bs), 4.3-3.4 (6H, m), 3.10 (1H, m), 2.59 (2H, m), 1.95 (2H, m), 1.44
- 15 (38) Methyl 2-(N'-t-butoxycarbonyl-N-2-carboxyethylhydrazino)-carbonyl hexahydropyridazine 3-carboxylate (522). Using a similar method to that described for 261 above, 522 was prepared, 92% as a white solid: mp. $146-148^{\circ}\text{C}$ (decomp); $\left[\alpha\right]_{\text{D}}^{22} + 27.8^{\circ}$ (c

(10H, m + s).

- 20 0.25, CH₂Cl₂); IR (KBr) 3346, 1740, 1710, 1626, 1497, 1290, 1250, 1206, 1179, 1159; ¹H NMR (CDCl₃) 87.60 (1H, bs), 7.5-5.5 (1H, vbs), 4.64 (1H, bs), 3.76 (5H, m + s), 3.00 (1H, m), 2.70 (3H, m), 2.16 (1H, m), 1.92 (1H, m), 1.56 (1H, m), 1.46 (11H, m + s). Anal. Calcd for
- 25 $C_{15}H_{26}N_4O_7$: C, 48.12; H, 7.00; N, 14.96. Found: C, 48.21; H, 6.96; N, 14.86. MS (ES^+) 373 $(M^- 1)$.

(4s) Methyl 7-t-butoxycarbonylamino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxylate (523).

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522 (7.15g, 19.1mmol) was dissolved in dichloromethane(100ml), containing dimethylformamide (0.5ml), and cooled to 0°C. Thionyl chloride (1.6ml, 2.61g, 22mmol) and N-ethyl morpholine (4.86ml, 440mg,

- 5 38.2mmol) were added and the mixture stirred for 2h. The organic mixture was washed with 2M sodium bisulphate (50ml), saturated sodium bicarbonate (50ml) and brine (50ml), dried (MgSO₄) and concentrated. The residues were triturated with ether to give 523 as a
- 10 white solid (5.73g, 84%): mp. 186-188°C (decomp); $\left\{\alpha\right\}_0^{22} + 65.3^\circ \text{ (c 0.25, CH}_2\text{Cl}_2\text{); IR (KBr) 3298, 2978,} \\ 1750, 1720, 1682, 1658, 1455, 1423, 1369, 1316, 1241, \\ 1212, 1160; ^1H \, \text{NMR (CDCl}_3\text{) } \delta 6.56 \, \text{(1H, s], 5.17 (1H, dd),} \\ 4.48 \, \text{(1H, bd), 3.81 (3H, m), 3.75 (3H, s), 2.83 (1H, s)}$
- 15 dt), 2.40 (1H, m), 2.28 (1H, m), 1.95 (1H, m), 1.67 (1H, m), 1.47 (9H, s). Anal. Calcd for $C_{15}H_{24}N_4O_6 * 1/6H_2O: C, 50.13; H, 6.82; N, 15.59. Found: C, 50.12; H, 6.71; N, 15.58. MS (ES^+) 357 (M^+ 1, 468), 301 (1008).$
- 20 (48) Methyl 7-amino-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4carboxylate (524), was synthesized from 523 via method used to prepare 518.

Compounds **262a-k** were synthesized via methods 25 used to prepare **211b-f**.

262a-k 263a-k

compound	R
262a 263a	Soi
262b 263b	
262c 263c	NHCO.
262d 263d	NHCO- OMe
262e 263e	Qi
262f 263f	
262g 263g	
262h 263h	i

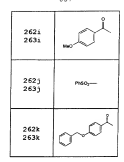
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20



- 25 (48) t-Butyl 6,10-dioxo-7-(2-naphthyl) sulfonamide-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxylate
 - (262a). 443mg (91%) of the title compound was obtained: mp. 56-7°C; $(\alpha)_{D}^{25}$ +76° (c 0.15, CH₂Cl₂); IR
- 30 (KBr) 3429, 2979, 1734, 1675, 1418, 1369, 1339, 1323, 1244, 1164, 665; ¹H NMR (CDCl₃) \(\delta \) 8.45 (1H, s), 8.00-7.59 (7H, m), 4.69-4.65 (1H, m), 4.25-4.12 (1H, m), 4.10-3.99 (1H, m), 3.73-3.55 (2H, m), 2.40-2.30 (1H, m), 1.99-1.91 (1H, m), 1.82-1.62 (2H, m), 1.48-1.46 (2H,
- 35 m), 1.37 (9H, s). Anal. Calcd for $C_{23}H_{28}N_4O_6S^{\bullet}H_2O^{\circ}$ C, 54.53; H, 5.97; N, 11.06. Found: C, 54.60; H, 5.73; N, 10.95. MS (ES $^+$) 489.
 - (4S) t-Butyl 6,10-dioxo-7-(3-methoxyphenylureido)-1,2,3,4,7,8,9,10-octahydro-6H-
- 40 pyridazino[1,2-a][1,2,4]triazepine-4-carboxylate
 (262c), 120mg (80%) of colourless foam was obtained:

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 $\begin{bmatrix} \alpha \end{bmatrix}_D^{22} + 22.6^{\circ} \ (c\ 0.1,\ CH_2Cl_2); \quad \text{IR} \ (KBr!\ 3316,\ 1732, \\ 1671,\ 1609,\ 1551,\ 1495,\ 1455,\ 1432,\ 1316,\ 1288,\ 1245, \\ 1218,\ 1158,\ 1122,\ 1023; \quad ^1_H\ NMR\ (CDCl_3)\ \delta 7.16\ (4H,\ m), \\ 6.79\ (1H,\ m)\ 6.60\ (1H,\ m),\ 5.11\ (1H,\ m),\ 4.59\ (1H,\ m), \\ 5\ 3.89\ (2H,\ m),\ 3.77\ (3H,\ s),\ 3.72\ (2H,\ m),\ 2.85\ (1H,\ m), \\ \end{bmatrix}$

(4S) t-Butyl 6,10-dioxo-7-(2-methoxyphenylureido)1,2,3,4,7,8,9,10-octahydro-6H-pyridazino
[1,2-a][1,2,4]triazepine-4-carboxylate (262d), (81%)
was obtained as colourless foam: [α]_D²² +3.7° (c 0.1,
10 CH₂Cl₂); IR (KBr) 3468, 3446, 3269, 1734, 1698, 1667,
1609, 1555, 1490, 1461, 1433, 1423, 1296, 1246, 1215,
1173, 1157, 1028, 756; ¹H NMR (CDCl₃) δ 8.23 (1H, m),
7.95 (1H, s), 6.95 (4H, m), 5.15 (1H, m), 4.60 (1H, m),
3.98-3.65 (4H, m), 3.89 (3H, s), 2.90 (1H, m), 2.48
15 (1H, m), 2.25 (1H, m), 2.05-1.65 (2H, m), 1.48 (9H, s).

(4S) t-Butyl 6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-7-phenylacetylamino-6H-

pyridazino[1,2-a][1,2,4]triazepine-4-carboxylate
(262e), was obtained as a white foamy solid (155mg,

- 20 53%): mp. 53-7°C; $\left[\alpha\right]_{D}^{22}$ +57.4° (c 0.1, CH₂Cl₂); IR (KBr) 3271, 2978, 1733, 1680, 1437, 1314, 1245, 1156; 1 H NMR (CDCl₃) δ 7.46 (1H, s), 7.42-7.20 (5H, m), 5.03 (1H, dd), 4.52-4.40 (1H, m), 3.96-3.70 (2H, m), 3.70-3.49 (1H, m), 3.63 (2H, s), 2.92-2.75 (1H, m), 2.43-
- 25 2.33 (1H, m), 2.33-2.15 (1H, m), 2.00-1.50 (3H, m), 1.45 (9H, s). Anal. Calcd for $C_{21}H_{28}N_4O_5 \cdot 0.25H_2O$: C, 59.91; H, 6.82; N, 13.31. Found: C, 60.19; H, 6.80; N, 13.30. MS (ES⁺) 418 (M⁺ + 2, 252), 417 (M⁺ + 1, 100), 362 (9), 361 (45).

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- (4s) t-Butyl 6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-7-(3-phenylureido)-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxylate (262f), was obtained as a white solid (273mq, 93%): mp. 102-6°C; [α]_n²² +7.5° (c 0.07,
- 5 CH₂Cl₂); IR (KBr) 3320, 2979, 1731, 1676, 1669, 1601, 1549, 1444, 1314, 1240, 1156; ¹H NMR (CDCl₃) δ 7.37-7.20 (6H, m), 7.08-6.98 (1H, m), 5.12 (1H, dd), 4.64-4.55 (1H, m), 4.02-3.78 (2H, m), 3.75-3.65 (1H, m), 2.94-2.75 (1H, m), 2.57-2.35 (1H, m), 2.35-2.20 (1H, m),
- 10 2.00-1.50 (3H, m), 1.48 (9H, s). Anal. Calcd for $C_{20}H_{27}N_{5}O_{5} \cdot 0.4H_{2}O$: C, 56.56; H, 6.60; N, 16.49. Found: C, 56.89; H, 6.58; N, 16.07. MS (ES⁺) 419 (M⁺ + 2, 24%), 418 (M⁺ + 1, 100), 363 (15), 362 (81), 242 (10).
 - (4S) t-Butyl 6,10-dioxo-7-(indole-2-carboxamido) -
- 15 1,2,3,4,7,8,9,10-octahydro-6H-pyridazino
 [1,2-a][1,2,4]triazepine-4-carboxylate (262g), (13g)
 was obtained as a white solid (298mg, 70%): mp. 13843°C; [α]₂²³ +69.8° (c 0.1, CH₂Cl₂); IR (KBr) 3282,
 2978, 1733, 1664, 1536, 1421, 1310, 1156, 748; ¹H NMR
- 20 (CDCl₃) δ 9.67 (1H, s), 9.53 (1H, s), 7.50 (1H, d), 7.307.15 (2H, m), 7.10-7.00 (1H, m), 6.93 (1H, s), 5.165.12 (1H, m), 4.60-4.50 (1H, m), 4.05-3.85 (2H, m),
 3.85-3.70 (1H, m), 3.05-2.90 (1H, m), 2.55-2.35 (1H,
 m), 2.35-2.20 (1H, m), 2.00-1.85 (1H, m), 1.85-1.50
- 25 (2H, m), 1.47 (9H, s). Anal. Calcd for $C_{22}H_{27}N_5O_5 \cdot 0.45H_2O$: C, 58.77; H, 6.26; N, 15.58. Found: C, 59.14; H, 6.24; N, 15.18. MS (ES⁺) 433 (M⁺ + 2, 26%), 442 (M⁺ + 1, 100), 387 (17), 386 (79), 285 (20), 229 (85), 211 (26), 185 (15), 183 (57), 139 (9).

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(4S) t-Butyl 7-[(4-acetamido)benzamido]-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]-triazepine-4-carboxylate (26Zh), was obtained as a white solid (325mg, 73%): mp. 209-12°C; [α]_D²⁴ +62.4°
5 (c 0.2, CH₂Cl₂); IR (KBr) 3513, 3269, 2980, 1731, 1680, 1653, 1599, 1531, 1314, 1158; ¹H NMR (CDCl₃) δ 9.40 (1H, s), 8.75 (1H, s), 7.72 (2H, d), 7.47 (2H, d), 5.15-5.05 (1H, m), 4.55-4.45 (1H, m), 4.05-3.70 (3H, m), 3.00-2.80 (1H, m), 2.45-2.35 (1H, m), 2.30-2.15 (1H, m), 1.48 (9H, s). Anal. Calcd for C₂₂H₂₉N₅O₆: C, 57.51; H, 6.36; N, 15.24. Found: C, 57.41; H, 6.38; N, 15.12. MS (ES^{*}) 461 (M^{*} + 2, 26%), 460 (M^{*} + 1, 100), 405 (12), 404 (55), 354 (7), 285 (23), 229 (52), 183 (22).

- 15 (4S) t-Butyl 6,10-dioxo-7-(4-methoxybenzoylamino) octahydro-6H-pyridazino[1,2-a][1,2,4]triazepinecarboxylate (262i), was obtained as a white glassy
 solid (76%): mp. 85-9°C; [α]_D²⁵ +66.4° (c 0.11,
 CH₂Cl₂); IR (KBr) 1732, 1668, 1607, 1502, 1440, 1312,
 20 1295, 1258, 1176, 1157, 1025; ¹H NMR (CDCl₃) δ 8.25 (1H,
 s), 7.77 (2H, m), 6.90 (2H, m), 5.11-5.07 (1H, m),
 4.55-4.48 (1H, m), 4.01-3.91 (2H, m), 3.86-3.78 (1H,
 m), 3.85 (3H, s), 2.98 (1H, m), 2.46-2.40 (1H, m),
 2.26-2.20 (1H, m), 2.05-1.80 (1H, m), 1.70-1.64 (2H,
 25 m), 1.48 (9H, s).
- (4s) t-Butyl 6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-7-phenylsulphonylamino-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxylate (262j), was obtained as a white crystalline solid

 30 (795): mp. 182-3°C (dec); [a]_n²² +92.1° (c 0.4, CH₂Cl₂);

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IR (KBr) 3283, 1732, 1684, 1448, 1430, 1404, 1369, 1338, 1306, 1285, 1242, 1169, 1091, 692; ^{1}H NMR (CDCl $_{3}$) δ 7.89 (2H, d, J = 7.4), 7.76 (1H, s), 7.64-7.49 (3H, m), 4.83 (1H, m), 4.35 (1H, brd, J = 13.0), 4.00 (1H, 5 m), 3.74-3.63 (2H, m), 2.39-2.26 (2H, m), 2.06 (1H, m), 1.50-1.41 (10H, m). Anal. Calcd for Cl $_{19}\text{H}_{26}\text{SN}_{4}\text{O}_{6}$: C, 52.04; H, 5.98 N, 12.78. Found: C, 52.11; H, 5.95; N, 12.71. MS (ES $^{+}$) 437 (M $^{+}$ - 1, 100%).

(3S) t-Butyl (7-(4-benzyloxyphenyl)carbonylamino-6,1010 dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino
[1,2-a][1,2,4]triazepine-4-carboxylate (262k), (83%)
was obtained: [α]_D²² +42.3°. (c 0.11, CH₂Cl₂);.IR (KBr)
3287, 2997, 2935, 1735, 1681, 1606, 1501, 1296, 1248,
1173,1155.

¹H NMR (CDCl₃) δ 9.23 (1H, s), 7.73 (2H, d),
15 7.38 (5H, m), 6.85 (2H, d), 5.08 (1H, m), 5.02 (2H, s),
4.48 (1H, bd), 4.15-3.65 (3H, m), 2.96 (1H, m), 2.452.10 (2H, m), 1.88 (1H, m), 1.63 (2H, m), 1.48 (9H, s).
M.S. (ES* 509 (M*-1).

Compounds 263a-k were synthesized via methods 20 used to prepare 212b-f.

(4S) 6,10-Dioxo-7-(2-naphthalenesulfonyl)amino-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263a), 348mg (94%) obtained as a white foamy solid:

25 mp. $[\alpha]_D^{21}$ +171° (c 0.056, CH_2Cl_2); IR (KBr) 3426, 3233, 2953, 1734, 1663, 1481, 1415, 1340, 1214, 1167, 1132, 1075, 668; 1H NMR ($CDCl_3$) δ 8.44 (1H, s), 8.00-7.60 (7H, m), 4.85-4.83 (1H, m), 4.25-4.00 (1H, m), 4.07-3.90 (1H, m), 3.70-3.46 (2H, m), 2.38-2.30 (1H, m), 2.12-

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2.01 (1H, m), 1.91-1.83 (1H, m), 1.46-1.26 (1H, m), 1.13-1.06 (1H, m), 0.90-0.77 (1H, m). MS (ES^{\dagger}) 431.

(4S) 7-(Benzo[b]thiophene-2-carbonyl)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

- 5 pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263b). 200mg (100%) was obtained as a white solid: mp. 155°C; $[\alpha]_D^{20}$ +13° (c 0.07, CH₂Cl₂); IR (KBr) 3431, 2935, 1734, 1663, 1531, 1435, 1292, 1177; 1_H NMR (CDCl₃) δ 9.73 (1H, bs), 7.73-7.27 (5H, m), 5.35-5.25
- 10 (1H, m), 4.56-4.48 (1H, m), 4.05-3.65 (3H, m), 3.12-3.00 (1H, m), 2.50-2.45 (1H, m), 2.30-2.20 (1H, m), 2.10-2.00 (1H, m), 1.75-1.61 (2H, m). MS (ES^{T}) 401.

(4S) 6,10-Dioxo-7-(3-methoxyphenylureido)-1,2,3,4,7,8,9,10-octahydro-6H-

- 15 pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263c), 216mg, (100+%) obtained as a colourless foam: $\left[\alpha\right]_D^{23}$ 32.5° (c 0.1, CH₂Cl₂); IR (KBr) 3326, 1730, 1661, 1610, 1555, 1495, 1431, 1314, 1288, 1217, 1175, 1161; 1 H NMR (CDCl₃) δ 7.87 (1H, s), 7.58 (1H, s), 7.19
- 20 (2H, m), 6.82 (1H, m), 6.62 (1H, m), 5.21 (1H, m), 4.55 (1H, m), 3.76 (3H, s), 4.0-3.65 (4H, m), 2.85 (1H, m), 2.35 (2H, m), 1.75 (1H, m), 1.71 (2H, m).

(4S) 6,10-Dioxo-7-(2-methoxyphenylureido)-1,2,3,4,7,8,9,10-octahydro-6H-

25 pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263d), (100+%) obtained as colourless foam: [\alpha]_0^{24} +11.7° (c 0.1, CH2Cl2); IR (KBr) 3394, 3325, 1666, 1603, 1543, 1490, 1463, 1438, 1329, 1311, 1292, 1249, 1214, 1176, 1119, 1024, 752; \$\frac{1}{14}\$ NMR (CDCl2) & 8.15 (18).

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m), 7.97 (2H, m), 7.15-6.84 (3H, m), 5.29 (1H, m), 4.62 (1H, m), 4.04-3.65 (4H, m), 3.89 (3H, s), 2.92 (1H, m), 2.50 (1H, m), 2.30 (1H, m), 2.10-1.75 (2H, m).

- (4S) 6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-7-
- 5 phenylacetyl-amino-6H-

pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263e), obtained as a white foamy solid (117mg, 98%): mp. 109-14 °C; [α] $_0^{24}$ +82.6° (c 0.06, CH₂Cl₂); IR (KBr) 3700-2250 (br), 3437, 3274, 2959, 1733, 1664, 1481,

- 10 1437, 1310, 1177; 1 H NMR (CDCl₃) δ 7.99 (1H, s), 7.40-7.15 (5H, m), 5.15-5.10 (1H, m), 5.25-4.70 (1H, bs), 4.50-4.35 (1H, m), 3.95-3.50 (3H, m), 3.61 (2H, s), 2.93-2.78 (1H, m), 2.40-2.20 (2H, m), 2.10-1.80 (1H, m), 1.80-1.60 (2H, m). Anal. Calcd for $C_{17}H_{20}N_{4}O_{5}\cdot H_{2}O_{5}\cdot H_{2}$
- 15 C, 53.96; H, 5.86; N, 14.81. Found: C, 54.12; H, 5.50; N, 14.68. MS (ES[†]) 360 (M+, 21%), 359 (M[†] 1, 100), 196 (14), 182 (14), 111 (7).
- (4S) 6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-7-(3-phenylureido)-6H-pyridazino[1,2-a][1,2,4]triazepine-4-20 carboxylic acid (263f), obtained as a white foamy solid (199mg, 92%): mp. 149-52°C; [α]_D²⁴ +92.0° (c 0.01, CH₃OH); IR (KBr) 3700-2300 (br), 3319, 2956, 1726, 1664, 1600, 1548, 1500, 1444, 1313, 1238, 755; ¹H NMR (D₆-DMSO)δ 8.90 (1H, s), 8.24 (1H, s), 7.42 (2H, d),
- 25 7.30-7.20 (2H, m), 7.00-6.90 (1H, m1, 4.98-4.92 (1H, m), 4.32-4.22 (1H, m), 3.80-3.55 (3H, m), 2.85-2.70 (1H, m), 2.30-2.20 (1H, m), 2.20-2.00 (1H, m), 1.90-1.35 (3H, m). Anal. Calcd for $C_{16}H_{19}N_{5}O_{5} \cdot 0.75H_{2}O_{5} \cdot c$, 51.26; H, 5.51; N, 18.68. Found: C, 51.11; H, 5.23; N,

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18.42. MS (ES^{\dagger}) 361 (M+, 208), 360 $(M^{\dagger} - 1, 100)$, 241 (11), 240 (89), 196 (15), 175 (29), 111 (12).

(4S) 6,10-Dioxo-7-(indole-2-carboxamido)-1,2,3,4,7,8,9,10-octahydro-6H-

- 5 pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263g), was obtained as a white solid (259mg, 92\$)mp. $248-51^{\circ}$ C; [α] $_{D}^{24}$ +94.0° (c 0.01, CH₃OH); IR (KBr) 3700-2300 (br) 3341, 2956, 1738, 1668, 1651, 1529, 1425, 1311, 1259, 751; 11 H NMR (D_{6} -DMSO) δ 13.29 (1H, bs),
- 10 11.72 (1H, s), 10.64 (1H, s), 7.65 (1H, d), 7.45 (1H, d), 7.26-7.15 (1H, m), 7.17 (1H, s), 7.10-7.00 (1H, m), 5.05-4.95 (1H, m), 4.40-4.25 (1H, m), 3.90-3.50 (3H, m), 2.88-2.75 (1H, m), 2.38-2.20 (1H, m), 2.20-2.00 (1H, m), 1.90-1.35 (3H). Anal. Calcd for
- 15 $C_{18}H_{19}N_{5}O_{5} \cdot 0.5H_{2}O$: C, 53.59; H, 5.25; N, 17.35. Found: C, 53.66; H, 4.88; N, 17.11. MS (ES^{+}) 385 $(M^{+}, 23\%)$, 384 $(M^{+} 1, 100)$, 298 (6), 253 (8), 227 (10), 199 (23), 196 (10), 173 (9), 126 (21).

(4S) 7-[(4-Acetamido)benzamido]-6,10-dioxo-

- 20 1,2,3,4,7,8,9,10-octahydro-6H
 - pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263h), was obtained as a white solid (282mg, 99°): mp. 210-5°C; $[\alpha]_D^{24}$ +74.5° (c 0.01, CH₃OH); IR (KBr) 3700-2300 (br) 3444, 3316, 2960, 1664, 1599, 1531, 1439,
- 25 1331, 1184; 1 H NMR (D_6 -DMS0) δ 13.30 (1H, bs), 10.50 (1H, s), 10.25 (1H, s), 7.80 (2H, d), 7.68 (2H, d), 5.00-4.90 (1H, m), 4.35-4.25 (1H, m), 3.90-3.40 (3H, m), 2.88-2.70 (1H, m), 2.35-2.25 (1H, m), 2.25-1.95 (1H, m), 2.08 (3H, s), 1.95-1.35 (3H, m). MS (ES †) 403

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 $(M+, 10%), 402 (M^{+} - 1, 100), 358 (10), 247 (10), 227$ (16), 219 (51), 198 (12), 184 (17).

- (4S) 6,10-Dioxo-7-(4-methoxybenzoylamino)-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-carboxylic acid
- 5 (263i), was obtained as a white glassy solid (approx 100%) used without purification: ¹H NMR (CDCl₂) 89.23 (1H, s), 7.72 (2H, d, J = 8.8), 6.81 (2H, d, J = 8.9), 5.22 (1H, m), 4.51 (1H, m), 3.97-3.72 (2H, m), 3.81 (3H, s), 3.03 (1H, m), 2.51-2.46 (1H, m), 2.31-2.25
- 10 (1H, m), 2.03 (1H, m), 1.72 (2H, m).
 - (4S) 6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-7phenylsulphonylamino-6H-

pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (2631), was obtained as a white solid (100%): mp. 73-

- 15 83°C (dec); $[\alpha]_{D}^{22} + 104.7^{\circ}$ (c 0.3, $CH_{2}Cl_{2}$); IR (KBr) 3600-2500 (br), 3208, 1734, 1666, 1481, 1448, 1416, 1338, 1311, 1214, 1171, 1091, 729, 689; ¹H NMR (CDCl₃) δ 7.87 (3H, m), 7.70-7.50 (3H, m), 7.16 (1H, brs), 4.99 $\{1H, m\}, 4.37 (1H, brd, J = 12.8), 3.92 (1H, m), 3.67$
- 20 $\{2H, m\}$, 2.36 $\{2H, m\}$, 2.13 $\{1H, brd, J = 12.2\}$, 1.56 (3H, m). Anal. Calcd for C15H18SN4O6*0.25CF3CO2H: C, 45.31; H, 4.48 N, 13.64. Found: C, 45.48; H, 4.71; N, 13.43. MS (ES⁺) 383 (MH⁺, 100%). Accurate mass calculated for $C_{15}H_{19}SN_4O_6$ (MH⁺): 383.1025. Found:
- 25 383.1007.
 - (4S) 7-(4-Benzyloxyphenyl)carbonylamino-6.10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (263k), (100%) obtained: mp. 130-142°C; IR (KBr) 3272,

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2945, 1738, 1650, 1611, 1501, 1445, 1309, 1255, 1171;

¹H NMR (CDCl₃) δ 9.35 (1H, s), 7.74 (2H, d), 7.38 (5H, m), 6.85 (2H, d), 5.40 (1H, bs), 5.19 (1H, s), 5.02 (2H, s), 4.49 (1H, d), 3.92 (2H, m), 3.68 (1H, m), 2.99 (1H, bs), 2.43 (1H, bs), 2.22 (1H, bs), 1.99 (1H, bs), 1.68 (2H, bs),

(4S) Methyl 6,10-dioxo-7-(3,4-methylenedioxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro10 6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxylate (5251), was synthesized via method used to prepare 211 to afford a white crystalline solid (3.35g, 83%): mp. 214-5°C; [α]_D²⁰ +75.2° (c 0.1, CH₂Cl₂); IR (KBr) 3272, 2955, 1747, 1664, 1610, 1485, 1443, 1265, 1040; ¹H NMR 15 (CDCl₃) δ 8.66 (1H, s), 7.32 (1H, dd), 7.23 (1H, d), 6.76 (1H, d), 6.02 (2H, s), 5.20 (1H, dd), 4.55-4.45 (1H, m), 4.03-3.70 (3H, m), 3.78 (3H, s), 3.05-2.88 (1H, m), 2.47-2.35 (1H, m), 2.35-2.20 (1H, m), 2.10-1.90 (1H, m), 1.85-1.50 (2H, m). Anal. Calcd for C₁gH₂ON₄O₇·0.5H₂O: C, 52.87; H, 5.06; N, 13.70. Found: C, 52.84; H, 5.00; N, 13.66. MS (ES*) 406 (M* + 2, 20%), 405 (M* + 1, 100), 391 (10), 162 (6), 148 (3),

(4*S*) 6,10-Dioxo-7-(3,4-methylenedioxybenzoylamino)-25 1,2,3,4,7,8,9,10-octahydro-6H-

105 (2).

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pyridazino[1,2-a][1,2,4]triazepine-4-carboxylic acid (2631). A suspension of 5251 (3.32g, 8.2mmol) in tetrahydrofuran (60ml) was treated with a solution of LiOH • H2O (0.69g, 16.4mmol, 2.0 equiv) in water (20ml). 5 The resulting mixture was stirred for 1h, concentrated and the residue dissolved in water (50ml). The solution was acidified using 2M. NaHSO4 and the product extracted with EtOAc (100ml and 50ml portions). The combined extract was washed once with brine (2 x 50ml), 10 dried (MgSO₄) and concentrated to afford 2631 as a white crystalline solid (2.87q, 90%): mp. 154-8°C; $[\alpha]_{\rm p}^{20}$ +85.6° (c 0.01, CH₂OH); IR (KBr) 3700-2300 (br), 3248, 2942, 1733, 1681, 1658, 1648, 1536, 1486, 1440, 1297, 1255, 1037; 1 H NMR (D₆-DMSO) δ 13.23 (1H, bs), 15 10.45 (1H, s), 7.45 (1H, d), 7.35 (1H, s), 7.03 (1H, d), 6.12 (2H, s), 5.00-4.93 (1H, m), 4.35-4.25 (1H, m), 3.90-3.40 (3H, m), 2.95-2.70 (1H, m), 2.40-2.25 (1H, m), 2.15-2.00 (1H, m), 1.91-1.40 (3H, m). Anal. Calcd for C₁₇H₁₈N₄O₇ • 0.8H₂O: C, 50.45; H, 4.88; N, 13.84.

20 Found: C, 50.80; H, 4.95; N, 13.36. MS (ES[†]) 390 (M[†], 19%), 389 (M[†] - 1, 100), 345 (9), 204 (31), 182 (27), 111 (12).

264a, c-1

265a, c, d, f 1015, 1018, 1027, 1052, 1056, 1075, 1095

2

compound	R ¹
264a 265a	So
264c 265c	ome H
264d 265d	H OM/e
264e 1095	Qi
264f 265f	
26 4 g 1075	
264h 1018	HO N
26 4 i 1052	Meo
264j 1027	50,

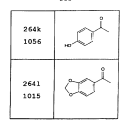
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[4S(2S,3S)] N-(2-Benzyloxy-5-oxo-tetrahydrofuran-3-yl)25 6,10-dioxo-7-(2-naphthalenesulfonyl)amino1,2,3,4,7,8,9,10-octahydro-6H-

pyridazino[1,2-a][1,2,4]triazepine-4-carboxamide
 (264a), was synthesized by a similar method as compound
213e to afford a white solid (240mg, 82%): IR (KBr)

330, 3066, 2947, 1789, 1750, 1691, 1454, 1417, 1368, 1298, 1262, 1235, 1193, 1118, 756, 696; 1 H NMR (D₆-DMSO) δ 8.59 (1H, d, J = 6.8), 8.48 (1H, s), 8.25-8.09 (3H, m), 7.85-7.75 (3H, m), 7.36 (5H, m), 5.39 (1H, m), 4.21 (2H, AB, J = 14.2), 4.53-4.49 (1H, m), 4.25-4.10

35 (2H, m), 3.65-3.44 (3H, m), 3.13-2.99 (1H, m), 2.43-2.16 (1H, m), 1.72-0.72 (7H, m). Anal. Calcd for $C_{30}H_{31}N_{5}O_{8}S: \text{ C, } 57.96; \text{ H, } 5.03; \text{ N, } 11.27. \text{ Found: C, } 57.28; \text{ H, } 5.14; \text{ N, } 10.48. \text{ MS } (ES^{+}) \text{ } 622.$

[4S(2S,3S)] N-(2-Benzyloxy-5-oxo-tetrahydrofuran-3-y1)-

40 6,10-dioxo-7-(3-methoxyphenylureido)-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-1carboxamide (264c), was prepared by a similar method as
213e, (55:) as a colourless foam: mp. 135-40°C; (α)_D²²
+51.6° (c 0.1, CH₂Cl₂); IR (KBr) 3314, 1790, 1664,

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1608, 1543, 1496, 1455, 1428, 1325, 1287, 1250, 1218, 1160, 1118, 1 H NMR (CDCl $_{3}$) δ 8.00 (1H, d, J = 7.1), 7.66 (1H, s), 7.55 (1H, s), 7.28 (5H, m), 7.14 (2H, m), 6.87 (1H, d, J = 7.4), 6.59 (1H, m), 5.42 (1H, s), 4.66 (5H, 5 m), 3.90-3.65 (4H, m), 3.73 (3H, s), 2.98 (2H, m), 2.38 (2H, m), 2.01-1.65 (3H, m).

 $\label{eq:condition} $$ \{4S(2S,3S)\} \ N^-(2-Benzyloxy-5-oxo-tetrahydrofuran-3-y1)-6,10-dioxo-7-(2-methoxyphenylureido)-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-1-$

- 10 **carboxamide (264d)**, was prepared by a similar method as **213e**, (72%) as colourless foam: $\left[\alpha\right]_D^{22}$ +21.4° (c 0.1, CH₂Cl₂); IR (KBr) 3302, 1791, 1689, 1678, 1664, 1602, 1536, 1489, 1461, 1437, 1420, 1249, 1119, 1023, 942, 751; 1 H NMR (CDCl₃) δ 8.07 (1H, d, J = 7.7), 7.82 (1H,
- 15 s), 7.68 (1H, d, J = 6.7), 7.49 (1H, s), 7.34 (5H, m), 6.96 (3H, m), 5.47 (1H, s), 4.82 (2H, d + m, J = 11.5), 4.63 (1H, d, J = 11.5), 4.49 (2H, m), 3.85 (4H, s + m), 3.68 (2H, m), 3.01 (2H, m), 2.46 (2H, m), 1.95 (3H, m), 1.57 (1H, m).
- 20 [4S(2RS,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-7phenylacetylamino-6H-

pyridazino[1,2-a][1,2,4]triazepine-4-carboxamide (264e)
was synthesized via a similar method as used to prepare

- 25 213e to afford a mixture of diastereomers (Syn:anti isomer ratio 9:1) as a white glassy solid (128mg, 78%): mp. 103-8°C; IR (KBr) 3419, 3302, 1793, 1664, 1535, 1421, 1327, 1256, 1123, 973; 1 H NMR (D₆-DMSO) δ 10.20 (0.9H, s), 9.35 (0.1H, s), 8.74 (0.1H, d), 8.49 (0.9H,
- 30 d), 7.36-7.15 (10H, m), 5.67 (0.9H, d), 5.44 (0.1H, s),

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4.85-4.75 (1H, m), 4.74-4.60 (1H, m), 4.77 and 4.63 (2H, dd), 4.30-4.10 (1H, m), 3.80-3.40 (3H, m), 3.43 (2H, s), 3.10-2.40 (3H, m), 2.25-2.15 (1H, m), 2.00-1.35 (4H, m). Anal. Calcd for $C_2gH_{31}N_5O_7 \cdot 0.5H_2O$: C, 5 60.21; H, 5.77; N, 12.53. Found: C, 60.38; H, 5.83; N, 12.13. MS (ES $^+$) 551 (M $^+$ + 2, 33%), 550 (M $^+$ + 1, 100), 480 (7), 343 (8), 279 (4).

[4s(2Rs,3s)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-y1)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-7-(3-

- phenylureido)-6H-pyridazino[1,2-a][1,2,4]triazepine-4carboxamide (264f), was prepared by a similar method as
 compound 213e to afford the pure syn-isomer as a white
 foamy solid (225mg, 82%): mp. 130-5°C; [α]_D²⁴ ±10.8° (c
 0.1, CH₂Cl₂); IR (KBr) 3316, 1791, 1688, 1676, 1664,
- 15 1601, 1536, 1445, 1314, 1242, 973; ¹H NMR (D₆-DMSO) δ 8.84 (1H, s), 8.49 (1H, d), 8.19 (1H, s), 7.45-7.18 (9H, m), 7.00-6.90 (1H, m), 5.68 (1H, d), 4.90-4.81 (1H, m), 4.75-4.60 (1H, m), 4.78 and 4.63 (2H, dd), 4.30-4.20 (1H, m), 3.75-3.55 (3H, m), 2.85-2.55 (3H,
- 20 m), 2.25-2.15 (1H, m), 2.00-1.35 (4H, m). Anal. Calcd for $C_2\gamma H_{30}N_6O_7 \cdot 0.5H_2O$: C, 57.95; H, 5.58; N, 15.02. Found: C, 58.12; H, 5.64; N, 14.81. MS (ES⁺) 552 (M⁺ 2, 30%), 551 (M⁺ + 1, 100), 362 (19), 299 (10), 279 (4).
- 25 [4\$(2\$,3\$)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)6,10-dioxo-7-(indole-2-carboxamido)-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4carboxamide (264g), was prepared by a similar method as
 compound 213e to afford the pure anti-isomer as a white
 30 solid (284mg, 80°): mp. 148-53°C; [α]n²⁴ -72.0° (c.0.1)

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CH₂Cl₂); IR (KBr) 3404, 3295, 1789, 1660, 1536, 1421, 1310, 1260, 1122, 749; 1 H NMR (D₆-DMSO) δ 11.72 (1H, s), 10.58 (1H, s), 8.73 (1H, d), 7.65 (1H, d), 7.58-7.27 (6H, m), 7.27-7.10 (1H, m), 7.17 (1H, s), 7.10-7.00 (1H, m), 5.46 (1H, s), 4.90-4.85 (1H, m), 4.77 and 4.68 (2H, dd), 4.35-4.25 (2H, m), 3.95-3.55 (3H, m), 3.09 (1H, dd), 2.95-2.80 (1H, m), 2.47-2.25 (2H, m), 2.10-1.35 (4H, m). MS (ES $^+$) 574 (M+, 35%), 573 (M $^+$ - 1, 100), 384 (16), 383 (69), 341 (23), 327 (12), 267 (13), 10 200 (22).

[4S(2RS, 3S)] 7-[(4-Acetamido)benzamido]-N-(2-benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4carboxamide (264h), was prepared by a similar method as 15 compound 213e to afford a mixture of diastereomers (Syn:anti isomer ratio 9:1) as a white solid (276mg, 70%): mp. 147-52°C; IR (KBr) 3444, 3304, 1793, 1665, 1602, 1531, 1505, 1423, 1294, 1264, 1181, 1123, 966; ¹H NMR (D₆-DMSO) δ 10.41 (1H, s), 10.22 (1H, s), 8.71 20 (0.1H, d), 8.48 (0.9H, d), 7.78 (2H, d), 7.67 (2H, d), 7.35-7.30 (5H, m), 5.68 (0.9H, d), 5.45 (0.1H, s), 4.88-4.80 (1H, m), 4.75-4.60 (1H, m), 4.77 and 4.63 (2H, dd), 4.30-4.20 (1H, m), 3.90-3.50 (3H, m), 3.10-2.50 (3H, m), 2.35-2.20 (1H, m), 2.07 (3H, s), 2.05-25 1.35 (4H, m). Anal. Calcd for C29H32N6O8+1H2O: C, 57.04; H, 5.61; N, 13.76. Found: C, 56.79; H, 5.50; N, 13.53. MS (ES^+) 594 $(M^+ + 2, 34\%)$, 593 $(M^+ + 1, 100)$. 387 (8), 386 (38), 358 (8), 162 (19).

[4S(2RS,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-yl)30 6,10-dioxo-7-(4-methoxybenzoylamino)-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxamide

(264i), was prepared by a similar method to that described for compound 213e to afford a white solid (70%): mp. 116-118°C; IR (KBr) 3315, 2951, 1793, 1664, 1607, 1502, 1258, 1177; ¹H NMR (CDCl₃) & 8.07 (1H, S),

- 5 7.77 (2H, d, J = 8.6), 7.35 (5H, m), 6.94 (2H, d, J = 8.5), 6.74 (1H), 4.89 (1H, d, J = 11.1), 4.74 (1H, m), 4.60 (1H, d, J = 11.0), 4.48, 4.41 (1H, 2m), 3.86 (3H, s), 3.79, 3.71-3.53 (3H, 2m), 2.87 (2H, m), 2.44 (1H, m), 2.18, 1.91, 1.68 (5H, 3m).
- 10 [4S(2S,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3yl)6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-7phenylsulphonylamino-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxamide
 (264j), was synthesized by a similar method as compound
- 15 **213e** to afford a foam (88%): $[\alpha]_D^{24}$ +74.2° (c 0.36, CH₂Cl₂); IR (KBr) 3332, 3235, 1793, 1664, 1537, 1448, 1416, 1337, 1169, 118, 1092, 940, 690; 1 H NMR (CDCl₃) δ 7.99 (1H, s), 7.88 (2H, d, J = 6.8), 7.64-7.48 (3H, m), 7.34 (5H, s), 7.13 (1H, d, J = 6.9), 5.39 (1H, s), 4.81
- 20 (2H, m), 4.62 (1H, d, J = 11.5), 4.48 (1H, m), 4.33 (1H, m), 3.85 (1H, m), 3.59 (2H, m), 3.03 (1H, dd, J = 7.6, 18.2), 2.49-2.28 (3H, m), 1.94-1.40 (4H, m). Anal. Calcd for $C_{26}H_{29}SN_{5}O_{8}$: C, 54.63; H, 5.11 N, 12.25. Found: C, 54.42; H,5.28; N, 11.62. MS (ES $^{+}$) 572 (MH $^{+}$,
- 25 100%). Accurate mass calculated for $C_{26}H_{30}SN_5O_8$ (MH *): 572.1815. Found: 572.1802.

[4S(2RS,3S)] 7-(4-Benzyloxyphenyl)carbonylamino-N-(2-benzyloxy-5-oxotetrahydrofuran-3-yl)-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

30 pyridazino[1,2-a][1,2,4]triazepine-4-carboxamide

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(264k), was prepared by the method used for 213e $(96\pm)$: IR (KBr) 3294, 2946, 1793, 1658, 1606, 1535, 1501, 1248, 1174, 1119. ¹H NMR (CDCl₃) δ 8.91 (1H, s), 7.85 (3H, m), 7.4 (10H, m), 7.02 (2H, d), 5.35 (1H, s), 5.10 (2H, s), 4.8-4.3 (5H, m), 4.00 (1H, bs), 3.78 (2H, m), 2.90 (2H, m), 2.5-1.5 (6H, m).

[45(2R5,3S)] N-(2-Benzyloxy-5-oxotetrahydrofuran-3-y1)-6,10-dioxo-7-(3,4-methylenedioxybenzoylamino)1,2,3,4,7,8,9,10-octahydro-6H-

- pyridazino[1,2-a][1,2,4]triazepine-4-carboxamide
 (2641), was prepared by a similar method as compound
 213e to afford a mixture of diastereomers (syn:anti
 isomer ratio 1:1) as a white solid (1.72g, 71%): mp.
 148-60°C; IR (KBr) 3314, 1780, 1677, 1658, 1651, 1550,
- 20 (2H, m), 3.65-3.43 (1H, m), 3.09 (0.5H, dd), 2.90-2.55 (1.5H, m), 2.45-2.10 (2H, m), 2.10-1.35 (4H, m), Anal. Calcd for $C_{28}H_{29}N_{5}O_{9} \cdot 0.2H_{2}O$: C, 57.67; E, 5.08; N, 12.01. Found: C, 58.01; H, 5.33; N, 11.51. MS (ES^T) 581 (M⁴ + 2, 33%), 580 (M+, 100), 374 (9), 373 (48),
- 25 345 (12), 261 (4), 239 (7), 149 (9).

[3S(4S)] 3-[6,10-Dioxo-7-(2-naphthalenesulfonyl)amino-1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4oxobutanoic acid (265a), was prepared by a similar 30 method as compound 265 to afford a white solid (37mg,

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17%): mp. 126-30°C (dec); $(\alpha)_D^{20} + 30^\circ$ (c 0.05, MeOH); IR (KBr) 3371, 2935, 1785, 1663, 1538, 1418, 1339, 1164, 669; 1 H NMR (CD₃OD) & 8.44 (1H, s), 8.06-7.50 (7H, m), 7.22 (1H, d, J = 8.4), 4.58-4.57 (1H, m), 4.46-4.42 (1H, m), 4.16-4.09 (2H, m), 3.85-3.50 (3H, m), 2.84-2.78 (1H, m), 2.64-2.51 (1H, m), 2.44-2.15 (2H, m), 1.81-0.89 (4H, m). Anal. Calcd for $C_{23}H_{25}N_{5}O_{6}S \cdot H_{2}O_{5}O_{5}S \cdot H_{2}O_{5}S \cdot H_{2}$

10 [3S(4S)] 3-[6,10-Dioxo-7-(3-methoxyphenylureido)1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4oxobutanoic acid (265c), was prepared by a similar
method as 265, (90%) as a colourless solid: mp. ~150°C

15 (decomp.); [α]_D²³ +94.8° (c 0.1, 20% MeOH/CH₂Cl₂); IR
(KBr) 3330, 1780, 1660, 1610, 1550, 1495, 1428, 1326,
1287, 1251, 1223, 1160; ¹H NMR (CD₃OD)δ7.16 (2H, m),
6.89 (1H, d, J = 7.8), 4.58 (1H, m), 4.37 (2H, m), 3.76
(6H, s + m), 2.95 (1H, m), 2.67 (1H, m), 2.33 (1H, m),
20 2.20-1.85 (3H, m), 1.66 (1H, m).

1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4-oxobutanoic acid (265d),
was prepared by a similar method as 265, (85%) as a
25 colourless solid: mp. ~176-85°C; (α)_D²³ +11.0° (c 0.1,
MeOH); IR (KBr) 3392, 3328, 1784w, 1665, 1603, 1537,
1490, 1462, 1437, 1337, 1290, 1290, 1217, 1177, 1119,
1023; ¹H NMR (CD₃OD) δ 8.02 (2H, m), 6.95 (4H, m), 5.05
(1H, m), 4.60 (2H, m), 3.92 (4H, s + m), 3.00 (2H, m),
30 2.68 (1H, m), 2.39 (1H, m), 2.00 (4H, m), 1.69 (1H, m).

[3S(4S)] 3-[6,10-Dioxo-7-(2-methoxyphenylureido)-

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[3S(4S)] 3-(6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-7-phenylacetylamino-6H-

pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido)-4oxobutanoic acid (1095), was prepared by a similar

- 5 method as compound **265** to afford a white solid (84mg, 90%): mp. 180-6°C; $\left[\alpha\right]_D^{22}$ +22.3° (c 0.065, CH₃OH); IR (KBr) 3700-2300 (br), 3287, 1664, 1536, 1425, 1261, 1181; 1 H NMR (CD₃OD) δ 7.35-7.20 (5H, m), 5.00-4.90 (1H, m), 4.60-4.50 (1H, m), 4.50-4.10 (2H, m), 3.90-3.50
- 10 (3H, m), 3.54 (2H, s), 3.00-2.80 (1H, m), 2.80-2.40 (2H, m), 2.35-2.20 (1H, m), 2.20-1.50 (4H, m), MS (ES *) 459 (M+ 24%), 458 (M * 1, 100), 358 (27), 175 (9), 149 (7), 137 (12). Accurate mass calculated for $C_{21}H_{26}N_{5}O_{7}$ (MH *): 460.1832. found: 460.1840.
- 15 [3S(4S)] 3-[6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-7-(3-phenylureido)-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4-oxobutanoic acid (265f), was prepared by a similar method as compound 265 to afford a white foamy solid (130mg, 88%): mp. 157-62°C; [α]_D²⁴ ·41.7°
 20 (c 0.1, CH₃OH); IR (KBr) 3700-2300 (br), 3325, 1782, 1663, 1547, 1443, 1315, 1242, 1181; ¹H NMR (CD₃OD) δ 7.40 (2H, dd), 7.35-7.20 (2H, m), 7.06-6.95 (1H, m), 5.05-4.95 (1H, m), 4.64-4.54 (1H, m), 4.50-4.35 (1H, m), 4.35-4.15 (1H, m), 3.90-3.69 (3H, m), 3.00-2.85 (1H, m), 2.80-2.45 (3H, m), 3.40-1.50 (4H, m) MS (ES¹)
 - 5 (1H, m), 2.80-2.45 (3H, m), 3.40-1.50 (4H, m). MS (ES^T 460 (M+, 24%), 459 (M^T 1, 100), 341 (9), 340 (54), 296 (6), 239 (9).

[3S(4S)] 3-[6,10-Dioxo-7-(indole-2-carboxamido)-1,2,3,4,7,8,9,10-octahydro-6H-

30 pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4-

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oxobutanoic acid (1075), was prepared by a similar method as compound 265 to afford a white solid (184mg, 83%): mp. 210-5°C; $\left(\alpha\right]_D^{24}$ +43.9° (c 0.1, CH₃OH), IR (KBr) 3700-2300 (br), 3309, 1660, 1537, 1423, 1311, 5 1262, 1184, 1_H NMR (CD₃OD) δ 7.61 (1H, d), 7.45 (1H, d), 7.28-7.15 (1H, m), 7.15-7.00 (1H, m), 7.13 (1H, s), 5.12-4.96 (1H, m), 4.62-4.55 (1H, m), 4.50-4.25 (2H, m), 4.00-3.69 (3H, m), 3.05-2.90 (1H, m), 2.80-2.30

(3H, m), 2.25-1.50 (4H, m). MS (ES^{+}) 484 (M+, 268), 10 483 $(M^{+} - 1, 100)$, 383 (25), 245 (12), 208 (11), 200 (21), 174 (31), 137 (18).

[3S(4S)] 3-{7-[(4-Acetamido)benzamido]-6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]-triazepine-4-carboxamido]-4-oxobutanoic acid (1018).

- 15 was prepared by a similar method as compound **265** to afford a white solid (177mg, 82%): mp. 235-40°C; $\left[\alpha\right]_{5}^{23}$ +27.3° (c 0.1, CH₃OH); IR (KBr) 3700-2300 (br), 3311, 2957, 1662, 1599, 1531, 1318, 1266, 1182; 1 H NMR (CD₃OD) δ 7.83 (2H, d), 7.69 (2H, d), 5.10-4.95 (1H, m),
- 20 4.64-4.55 (1H, m), 4.50-4.35 (1H, m), 4.32-4.22 (1H, m), 4.00-3.65 (3H, m), 3.05-2.90 (1H, m), 2.80-2.30 (3H, m), 2.15 (3H, s), 2.15-1.50 (4H, m). Anal. Calcd for $C_{22}H_{26}N_{6}O_{8} \cdot 1.5H_{2}O$: C, 49.90; H, 5.52; N, 15.87. Found: C, 50.21; H, 5.41; N, 15.49. MS (ES †) 502 (M+,
- 25 28%), 501 (M⁴ 1, 100), 401 (8), 218 (4), 119 (2), 116 (5), 113 (16).

[3S(4S)] 3-[6,10-Dioxo-7-(4-methoxybenzoylamino)octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4carboxamido]-4-oxobutanoic acid (1052), was synthesized
30 via method used to prepare 265 to afford a white solid

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(0.194g, 100%): mp. 138-142°C; $\left(\alpha\right)_{D}^{20}$ +36.3° (c 0.19, CH₃OH); IR (KBr) 3434-2962, 1782, 1660, 1607, 1537, 1504, 1441, 1424, 1313, 1293, 1258, 1177; ¹H NMR (CD₃OD) δ 7.11 (2H, d, J = 8.8), 6.90 (2H, d, J = 8.9), 4.46 (1H, m), 4.34, 4.28 (1H, 2m), 4.15 (1H, m), 3.75 (3H, s), 3.75, 3.70 (3H, m), 2.88, 2.49, 2.28, 2.23, 2.00, 1.86, 1.79, 1.58 (8H, m).

[3S(4S)] 3-(6,10-Dioxo-1,2,3,4,7,8,9,10-octahydro-7-phenylsulphonylamino-6H-

- 10 pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido)-4oxobutanoic acid (1027), was synthesized by a similar
 method as compound 265 to afford a white foam (88%):
 [α]_D²⁴ +22.6° (c 0.17, MeOH); IR (KBr) 3349, 1789,
 1663, 1537, 1448, 1337, 1169, 1092, 690; ¹H NMR (CD₃OD)
- 15 δ 7.82 (2H, d, J = 7.8), 7.57 (3H, m), 4.74 (1H, m), 4.47 (1H, m), 4.24-4.10 (2H, m), 3.72-3.47 (4H, m), 2.62-2.48 (3H, m), 2.20 (1H, m), 1.94-1.35 (3H, m). MS (ES⁺) 480 (M⁺ 1, 100%). Accurate mass calculated for $C_{19}H_{24}Sn_{5}O_{8}$ (MH⁺): 482.1346. Found: 482.1325.
- 20 [3S(4S)] 3-[6,10-Dioxo-7-(4-hydroxybenzoylamino)1,2,3,4,7,8,9,10-octahydro-6Hpyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4oxobutanoic acid (1056), was prepared by the method
 used for 265 (95%): mp. >300°C; IR (KBr) 3392, 1660,
 25 1610, 1507, 1442, 1280, 1171, 1149, 1133.

 H NMR
 (CD₃OD) δ7.74 (2H, d J = 8.7), 6.84 (2H, d J = 8.7) 4.58
 (1H, m), 4.41 (1H, bd, J = 12.6), 4.28 (1H, m), 3.85
 (3H, m), 2.98 (1H, m), 2.8-2.3 (3H, m), 2.3-1.6 (4H,

m).

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[35(45)] 3-[6,10-Dioxo-7-(3,4-

methylenedioxybenzoylamino) -1,2,3,4,7,8,9,10-octahydro-6H-pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4-oxobutanoic acid (1015), was prepared by a similar method as used for 265 to afford a white solid (142mg, 58%): mp. 170-5°C; [α]_D²⁵ +32.7° (c 0.1, CH₃OH); IR (KBr) 3700-2500 (br), 3325, 2969, 1784, 1662, 1485, 1440, 1292, 1258, 1037; ¹H NNR (CD₃OD) &7.45 (1H, dd), 7.32 (1H, d), 6.90 (1H, d), 6.05 (2H, s), 5.10-4.90 (1H, m), 4.62-4.54 (1H, m), 4.45-4.35 (1H, m), 4.33-4.22 (1H, m), 3.95-3.65 (3H, m), 3.05-2.90 (1H, m), 2.80-2.30 (3H, m), 2.20-1.50 (4H, m).

[3S(4S)] t-Butyl 3-[7-(benzo[b]thiophene-2-carbonyl)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-15 6H-pyridazino[1,2-a][1,2,4]triazepine]-4-oxobutanoate semicarbazone (526), was prepared by a similar method as used for 502 to afford a glassy solid: [α]_D²⁰ +34° (c 0.13, CH₂Cl₂); IR (KBr) 3437, 2929, 1670, 1530, 1428, 1288, 1156; ¹H NMR (CDCl₃) δ10.0 (1H, bs), 9.74 (1H, bs), 7.93 (1H, s), 7.80-7.60 (2H, m), 7.40-7.18 (3H, m), 6.15-5.30 (2H, bs), 5.00-4.85 (2H, m), 4.50-4.25 (1H, m), 3.95-3.75 (3H, m), 3.12-2.78 (2H, m), 2.73-1.60 (7H, m), 1.36 (9H, s). Anal. Calcd for

- 680 -

 $C_{27}H_{34}N_{8}O_{7}S$: C, 52.76; H, 5.58; N, 18.23. Found: C, 52.25; H, 5.74; N, 16.30. MS (ES^{+}) 615.

[3s(4s)] 3-[7-(Benzo[b]thiophene-2-carbonyl)amino-6,10-dioxo-1,2,3,4,7,8,9,10-octahydro-6H-

- 5 pyridazino[1,2-a][1,2,4]triazepine-4-carboxamido]-4oxobutanoic acid (1053), was prepared by a similar
 method as used for 214 to afford a white solid (106mg,
 73%): [\alpha]_0^2 +22\alpha (c 0.10, MeOH); IR (KBr) 3428, 2944,
 1733, 1652, 1532, 1433, 1337, 1288, 1186; \frac{1}{1}H NMR
- 10 (CD₃OD) δ 7.95 (1H, s), 7.90-7.85 (2H, m), 7.43-7.35 (2H, m), 4.98 (1H, m), 4.65-4.52 (1H, m), 4.40-4.20 (2H, m), 3.85-3.70 (3H, m), 3.30-3.25 (3H, m), 3.03-2.85 (1H, m), 2.70-2.31 (3H, m), 2.10-1.55 (4H, m). MS (ES⁺) 500 (as methyl acetal of the aldehyde).

528

15

[4S(2RS,3S)] 6,10-Dioxo-N-(2-ethoxy-5-oxotetrahydrofuran-3-yl)-7-(3,4-methylenedioxybenzoylamino)-1,2,3,4,7,8,9,10-octahydro-6H-pyridazıno[1,2-a][1,2,4]triazepine-4-carboxamide

20 (528), was prepared by a similar method as compound

213e to afford a mixture of diastereomers (Syn: antiisomer ratio 1:1) as a creamy white foamy solid (1.05a,

- 681 -

58%): mp. 124-32°C; IR (KBr) 3312, 2979, 1790, 1664, 1610, 1532, 1485, 1285, 1120, 1037, 932; 1 H NMR (D₆-DMSO) δ 10.39 (1H, s), 8.71 (0.5H, d), 8.43 (0.5H, d), 7.45 (1H, d), 7.36 (1H, s), 7.04 (1H, d), 6.12 (2H, s), 5.58 (0.5H, d), 5.34 (0.5H, s), 4.95-4.85 (1H, m), 4.70-4.52 (0.5H, m), 4.35-4.10 (1.5H, m), 3.95-3.50 (5H, m), 3.03 (0.5H, dd), 2.90-2.55 (1.5H, m), 2.46-2.20 (2H, m), 2.10-2.40 (4H, m), 1.16-1.13 (3H, 2 x t). Anal. Calcd for $C_{29}H_{27}N_{5}O_{9} \cdot 0.6H_{2}O$: C, 52.29; H, 5.38; N, 10 13.26. Found: C, 52.53; H, 5.35; N, 12.78. MS (ES[†]) 519 (M[†] + 2, 278), 518 (M[†] + 1, 100), 472 (7), 374

Example 31

Compounds 640, 642, 645, 650, 653, 655, 656,

15 662, 668, 669, 670, 671, 677, 678, 681, 682, 683, 684, 686, 688a, 688b, 6891, 689b, 690a, 690b, 691a, 691b, 695a, 695b, 695c, 692a, 692b, 693 and 694 were prepared as follows.

(12), 373 (53), 345 (14), 149 (12).

- 682 -

(35)-2-0xo-3-amino-5-methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid methyl ester (638), was synthesized from 600a by methods similar to those used for making 602m from 600a to afford 2.4g of 638 as 5 a white solid.

(3S)-2-0xo-3-(2-naphthylmethylene)amino-5methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepinel-acetic acid methyl ester (639). To a solution of 638
(630 mg, 1.76 mmol) and 2-naphthylmethyl bromide (428)
mg, i.94 mmol) in CH₃CN was added K₂CO₃ (608 mg, 4.4
mmol). The resulting mixture was stirred at ambient
temperature. After 18 hours, the reaction mixture was
diluted with CH₂Cl₂, washed with water then brine,
dried over Na₂SO₄ then concentrated in vacuo. Flash
thromatography (SiO₂, 0 to 20% EtOAc/CH₂Cl₂) afforded
450mg of 639.

(3S)-3-[(3S)-2-0xo-3-(2-naphthylmethylene)amino-5-methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (640), was synthesized by methods used to make 605v from 602v to afford 205 mg of 640 as a white solid, ¹H NMR (CDCl₃) & 2.4-2.55(m, 1H), 2.65-2.8(m, 1H), 3.2(s, 3H), 3.72-3.78(m, 1H), 3.85-4.0(m, 2H), 4.22-4.28(d, 1H), 4.26-4.5(m, 4H), 4.58-4.75(m, 1H), 4.78-4.85(m, 1H), 5.0-5.08(t, 1H), 7.35-7.65(m, 7H), 7.85-8.02(m, 4H).

(35) -3-[(35) -2-0xo-3-benzoylformylamino-5-methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (642), was synthesized from 638 by similar methods used to make 605m to afford 5 213 mg of 642, ¹H NMR (CD₃OD) δ 2.5(m, 1H), 2.68 (ddd, 1H), 3.25(s, 2H), 3.3(s, 3H), 3.78(m, 2H), 4.0(d, 1H), 4.3(m, 1H), 4.6(m, 2H), 4.85(br. s, 2H), 7.08-7.22(m, 2H), 7.35(m, 1H), 7.4-7.65(m, 4H), 7.7(dd, 1H), 8.1(dd, 2H), 7.1(dd, 2H), 8.1(dd, 2H), 8.1(

1H).

10 2-Acetamido-acetyl chloride (643). To a suspension of N-acetyl glycine (200 mg, 1.7 mmol) in CH₂Cl₂ (2.5 mLs) containing DMF (0.005 mLs) was added oxalyl chloride

- 684 -

(0.450 mLs, 5.1 mmol). After stirring 30 minutes at ambient temperature, the mixture was concentrated to afford 643 as a crude product.

(35)-2-0xo-3-(1-naphthoy1) amino-5-(2-acetamido) acetyl5 2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid
benzyl ester (644), was synthesized from 600b by
methods used to make 602d from 600b using 643 to afford
112 mg of 644.

(3S)-3-[(3S)-2-Oxo-3-(1-naphthoy1) amino-5-(210 acetamido) acetyl-2,3,4,5-tetrahydro-1H-1,5benzodiazepine-1-acetylamino]4-oxo-butyric acid (645),
was synthesized from 644 by methods used to make 605d
from 602d to afford 43 mg of 645 as a white solid, ¹E
NMR (CD₃OD) δ 1.95(s, 3H), 2.4(m, 1H), 2.65(m, 1H),

15 3.4(s, 1H), 3.55(m, 1H), 3.85(m, 1H), 4.05(d, 1H), 4.3(m, 1H), 4.4-4.6(m, 2H), 5.0(m, 1H), 7.4-7.7(m, 6H), 7.85-8.0(m, 2H).

- 2-(N-Methyl, N-fluorenylmethoxycarbonyl)aminoacetyl
 chloride (646), was prepared from N-Fmoc-sarcosine by
 method used to make 643 to afford 646 as a crude
 product.
- 5 (3S)-2-Oxo-3-(1-naphthoyl)amino-5-[2-(N-methyl, N-fluorenylmethoxycarbonyl) amino]acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid benzyl ester (647), was synthesized from 600b by methods used to synthesize 602d from 600b, using 646 to afford 481 mg of 647.
 - (3S)-3-[(3S)-2-0xo-3-(1-naphthoyl)amino-5-[2-(N-methyl, N-fluorenylmethoxycarbonyl)amino]acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid tert-butyl ester semicarbazone (648), was

- 686 -

synthesized from 647 by methods used to prepare 604d from 602d to afford 409 mg of 648.

(3S)-3-[(3S)-2-0xo-3-(1-naphthoy1)amino-5-(2-methy1) amino) acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-5 1-acetylamino]4-oxo-butyric acid tert-buty1 ester

A solution of **648** (409 mg, 0.465 mmol) in MeCN:Et₂NH (4:1, v/v) was stirred at ambient temperature. After 45 minutes, the reaction mixture was concentrated in 10 vacuo. Flash chromatography (SiO₂, 5% to 20% MeOH in CH₂Cl₂) afforded 241 mg of **649**.

semicarbazone (649).

(38) -3-[(38) -2-0xo-3-(1-naphthoy1) amino-5-(2-methy1 amino) acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino)4-oxo-butyric acid (650), was synthesized 15 from 649 by methods used to prepare 605d from 604 to afford 179 mg of 650 as a white solid, ¹H NMR (CD₃OD) δ 2.4-2.6(m, 2H), 2.7(s, 3H), 3.5(q, 1H), 3.8 (m, 2H), 4.2-4.4 (m, 2H), 4.3-4.45(m, 1H), 5.0-5.1(m, 2H), 7.4-7.7(m, 6H), 7.85-7.9(m, 2H), 8.2 (m, 1H).

20 (3S)-2-0xo-3-(1-naphthoyl)amino-5-formyl-2,3,4,5tetrahydro-1H-1,5-benzodiazepine-1-acetic acid benzyl ester (652), was synthesized from 600b by methods similar to those used to make 602n from 600b, using the

reagent obtained from reacting DMF with 3 equiv. of oxalyl chloride in a ${\rm CH_2Cl_2}$ solution as ${\rm R}^3{\rm X}$, to afford 404 mg of 652.

(3S) -3 - [(3S) -2 -0xo -3 - (1-naphthoy1) amino -5 -formy1 -

5 2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (653), was synthesized from 652 by methods used to prepare 605d from 602d to afford 84 mg of 653 as a white solid, ¹H NMR (CD₃OD) δ 2.3 (m, 1H), 2.55 (dd, 1H), 3.75 (br. s, 1H), 4.25-4.6 (m 10 5H), 5.15 (m, 1H), 7.2-7.45 (m, 6H), 7.8-7.9 (dd, 3H),

8.1(s, 1H), 8.2(m, 2H).

(3S)-2-Oxo-3-(3,5-dichloro-4-hydroxybenzoyl)amino-5acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1acetic acid (654), was synthesized from 600b using 15 methods similar to those used for preparing 603d from 600b to afford 775 mg of 654.

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(3S)-2-Oxo-3-(3,5-dichloro-4-hydroxybenzoy1) amino-5-acety1-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-y1]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (655), was synthesized from 654 using the method used to prepare 213e to afford 304 mg of 655, ¹H NMR (CD₃OD) & 2.4(d, 1H), 2.6-2.75(m, 2H), 3.0(m, 1H), 3.45(m, 1H), 3.8(d, 1H), 4.0(t, 2H), 4.4(m, 2H), 4.5-4.55(m, 2H), 7.2-7.45(m, 4H), 7.85(s, 2H).

(3S)-3-[(3S)-2-0xo-3-(3,5-dichloro, 4-

- 10 hydroxybenzoyl)amino-5-acetyl-2,3,4,5-tetrahydro-1H1,5-benzodiazepine-1-acetylamino|4-oxo-butyric acid
 (656), was synthesized from 655 using a method similar
 to that used to prepare 2002 from 2001 to afford 136 mg
 of 656 as a white solid, ¹H NMR (CD₃OD) & 1.85(s, 3H),
- 15 2.5(m, 1H), 2.65(m, 1H), 3.7(m, 1H), 4.3(m, 1H), 4.55(m, 2H), 7.4-7.6(m, 4H), 7.85(s, 2H).

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2-(Fluorenylmethoxycarbonyl)hydroxyacetic acid benzyl ester (657). To a solution of benzyl glycolate (6.0 g, 36.1 mmol) in CH₂Cl₂, cooled via ice-water bath, was added fluorenylmethoxy chloroformate (14 g, 1.5 equiv.) 5 then diisopropylethylamine (9 mLs, 1.5 equiv.). After 1 hour, reaction mixture was poured into a saturated aqueaous solution of ammonium chloride and extracted with CH₂Cl₂, dried over Na₂SO₄ then concentrated in vacuo. The product was triturated from MeOH to obtain 10 2.2 q of 657 as a first crop of white solid.

- 2-(Fluorenylmethoxycarbonate) acetic acid (658). To a solution of 657 (2.2 g, 5.93 mmol) in tetrahydrofuran, was added 5% Pd/C (220 mg). The resulting suspension was vigorously stirred under hydrogen atmosphere.
- 15 After 90 min, the reaction mixture was filterred through Celite. The filtrate was poured into saturated aqueous NaHCO3 and washed twice with EtOAc. The aqueous layer was then acidified and the product extracted twice with CH2Cl2, dried over Na2SO4 and 20 concentrated in vacuo to afford 1.46 g (88%) of 658 as a white solid.
 - 2-(Fluorenylmethoxycarbonate) acetyl chloride (659), was prepared from 658 by the method used to prepare 643 to afford 659 as a crude product.
- 25 (3S) -3-[(3S) -2-Oxo-3-(3,5-dichloro-4hydroxybenzoyl) amino-5-(2fluorenylmethoxycarbonate) acetyl-2,3,4,5-tetrahydro-1H1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid
 tert-butyl ester semicarbazone (660), was synthesized

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from 600b, using 659, by methods used to prepare 604d from 600b to afford 453 mg of 660.

(3S) -3-[(3S) -2-0xo-3-(3,5-dichloro-4-

hydroxybenzovl)amino-5-(2-hydroxy)acetyl-2,3,4,5-

- 5 tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxobutvric acid tert-butvl ester semicarbazone (661). A solution of 660 (423 mg) in MeOH:EtaNH (1:1, v/v) was stirred at ambient temperature. After 10 minutes, the reaction mixture was concentrated in vacuo to a small 10 volume. Precipitation by the addition of ether
 - afforded 230 mg of 661.

(3S) -3 - [(3S) -2 - 0xo -3 - (3,5 - dichloro -4 -

hydroxybenzoyl)amino-5-(2-hydroxy) acetyl-2,3,4,5-

tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-

- 15 butyric acid (662), was synthesized from 661 by the methods used to prepare 605d from 604 to afford 37 mg of 662 as a white solid, $^{1}\text{H NMR (CD}_{3}\text{OD)}$ δ 2.45(m, 1H), 2.7(m, 1H), 3.75(m, 1H), 3.9(d, 1H), 4.15(d, 1H), 4.35(m, 1H), 4.5(t, 2H), 4.7(dd, 1H), 7.4-7.6(m, 4H),
- 20 7.85(s. 2H).

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2-(Triisopropylsilyloxy)acetic acid benzyl ester (663).

To a solution of benzyl glycolate (46.91g, 0.282 mol) and diisopropylethylamine (74 mLs, 0.423 mol) in CH₂Cl₂, cooled via water bath, was added a solution of TIPSOTF (95 g, 0.31 mol) in CH₂Cl₂. The resulting mixture was allowed to warm to ambient temperature then poured into water, washed twice with 10% aqueous NaHSO₄, dried over Na₂SO₄ and concentrated *in vacuo*. Flash chromatography (SiO₂, 0 to 5% EtOAc in hexanes)

10 afforded 71.6 g of 663.

- 2-(Trisopropylsilyloxy)acetic acid (664). To a solution of 663 (0.4 g, 1.2 mmol) in EtOAc was added 10% Pd/C (33 mg). The resulting suspension was stirred under hydrogen atmosphere. After 15 hours, the reaction mixture was filterred through Celite and the filtrate concentrated in vacuo to afford 0.29 g of an oil. To a solution of this oil in 1,4-dioxane was added NaHCO3 (0.5M, 2.4 mLs). The resulting solution was concentrated in vacuo from toluene to afford 664 as
 - 2-(Triisopropylsilyloxy)acetyl chloride (665), was synthesized from 664 by a method similar that used to prepare 643 to afford 665 as a crude product.

(3S) -3-[(3S) -2-Oxo-3-benzoylamino-5-(2-

20 a waxy solid.

25 triisopropylsilyloxy)acetyl-2,3,4,5-tetrahydro-1H-1,5benzodiazepine-1-acetylamino]4-oxo-butyric acid tertbutyl ester semicarbazone (666), was synthesized from
600b, using 665, by methods used to prepare 604d from
600b to afford 131 mg of 666.

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(38)-3-[(38)-2-0xo-3-benzoylamino-5-(2-hydroxy)acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid tert-butyl ester semicarbazone (667). To a solution of 666 (131 mg, 0.17 mmol) in tetrahydrofuran, cooled via ice-water bath, was added tetrabutylammonium fluoride (1M, 0.190 mL). After 2 hours the reaction mixture was poured into water, extracted twice with EtOAc, dried over MgSO₄ and concentrated in vacuo to afford 63 mg of 667 as a white

(3s)-3-[(3s)-2-0xo-3-benzoylamino-5-(2-hydroxy) acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (668), was synthesized from 667 by the methods used to prepare 605d from 604d to afford 48 mg of 668 as a white solid, ¹H NMR (CD₃OD) δ 2.45 (m, 1H), 2.67 (dddd, 1H), 3.78 (d, 1H), 3.85 (br. m, 1H), 4.05 (d, 1H), 4.28 (m, 1H), 4.5 (m, 2H), 4.65 (m, 1H), 4.95 (br. s, 2H), 7.4-7.5 (m, 4H), 7.52-7.65 (m, 3H), 7.88 (d, 2H).

20 (3s)-3-[(3s)-2-0xo-3-(3,5-dichloro-4-methoxybenzoyl)amino-5-acetyl-2,3,4,5-tetrahydro-1H1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid
(669), was synthesized from 600b by the methods used to
prepare 605d from 600b to afford 63 mg cf 669 as a

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white solid, 1H NMR (CD₃OD) δ 1.9(s, 3H), 2.4-2.7(m, 2H), 3.6-3.7(m, 2H), 3.9(s, 3H), 4.2-4.4(m, 2H), 4.4-4.6(m, 3H), 7.4-7.8(m, 4H), 7.9(s, 2H).

(3S)-2-Oxo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-acetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (670), was synthesized from 600b by the methods used to prepare 655 from 600b to afford 218 mg of 670 as a white solid, ¹H NMR (CD₃OD) δ 1.7, 1.75(2s, 10 3H), 2.15, 2.2(2s, 6H), 2.4-2.5(m, 1H), 2.6-2.75(m, 1H), 3.65-3.75(m, 2H), 4.2-4.3(m, 2H), 4.45-4.6(m, 3H), 7.35-7.6(m, 4H), 7.5(s, 2H).

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(3S)-2-0xo-3-tert-butoxycarbonylamino-5-(2-triisopropylsilyloxy)acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid benzylester (672), was synthesized from 600b by method 1 used to prepare 602n

- 5 from 600b using 665 to afford 1.08 g of 672.
- (3S)-2-0xo-3-amino-5-(2-triisopropylsilyloxy)acetyl2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid
 benzylester (673). To a solution of 672 (1.08 g, 1.69
 mmol) in CH₂Cl₂ was added 2,6-lutadine (0.8 mL) then
 10 TMSOTf (1 mL, 5.1 mmol). After 1 hour, the reaction
 mixture was poured into NaHCO₃ and extracted with
 CH₂Cl₂, dried over MgSO₄ and concentrated in vacuo to a
 small volume that was used directly for the next
 reaction.
- 15 (3S)-2-Oxo-3-(1,6-dimethoxybenzoyl formyl)amino-5-(2-triisopropylsilyloxy)acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid benzylester (674), was synthesized from 673 by the method used to prepare 602b to afford 0.91 g of 674.
- 20 (3s)-2-0xo-3-(1,6-dimethoxybenzoyl formyl)amino-5-(2-triisopropylsilyloxy)acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid (675). A solution of 674 (0.365 g, 0.5 mmol) in MeOH was stirred with 1N NaOH (1.2 mL, 1.2 mmol). After 16 hours the reaction.
- 25 mixture was concentrated in vacuo then dissolved in water and washed twice with ether. The aqueous layer was acidified with $1\underline{N}$ HCl and the product extracted with EtOAc, dried over MgSO $_4$ and concnetrated in vacuo to afford 337 mg of 675 as a solid.

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(3S)-2-0xo-3-(1,6-dimethoxybenzoylformyl) amino-5-(2-triisopropylsilyloxy)acetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (676), was synthesized from 675 by the method used to prepare 213e to afford 166 mg of 676 as a white solid.

(3S)-2-Oxo-3-(1,6-dimethoxybenzoylformyl)amino-5-(2-hydroxy)acetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5
10 benzodiazepine-1-acetamide (677). A solution of TBAF (6 mL, 3 mmol) in HOAc (0.46 mL, 8 mmol) was added to 676 (0.213 g, 0.256 mmol). After 16 hours the reaction mixture was poured into EtOAc and washed twice with NaHCO₃, once with brine then dried over MgSO₄ and 15 concnetrated in vacuo to afford 139 mg of 677 as a solid, ¹H NMR (CDCl₃) δ 2.4(d, 1H), 2.5(dd, 1H), 2.8(dd, 1H), 2.92(dd, 1H), 3.15(m, 2H), 3.55-3.65(m, 2H), 3.72(s, 6H), 3.92(m, 1H), 4.05(m, 1H), 4.3(m, 1H), 4.42(d, 1H), 4.6(dd, 1H), 4.65-4.8(m, 2H), 4.88(d, 1H), 2.55(d, 1H), 6.55(m, 2H), 6.75(d, 1H), 7.25-7.55(m, 6H), 7.75(m, 2H).

5-(2-hydroxy)acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (678).
25 was synthesized by the method used to prepare 667 from 666 to afford 54 mg of 678 as a white solid, $^1\mathrm{H}$ NMR (CD₃OD) δ 2.45(m, 1H), 2.7(m, 1H), 3.5(m, 2H), 3.75(br. s, 6H), 4.05(d, 1H), 4.3(m, 1H), 4.51-4.6(m, 2H), 4.8(br. m, 2H), 6.7(d, 2H), 7.4-7.5(br. m, 3H), 7.6-30 7.65(br. m, 2H).

(3S) -3-[(3S) -2-0xo-3-(3,5-dimethoxybenzoylformyl) amino-

(35)-2-0xo-3-benzoylformylamino-5-(2-hydroxy)acetyl-N-(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (680), was synthesized from 600b by the methods used to prepare

- 5 677 from 600b to afford 140 mg of 680 as a white solid,

 ¹H NMR (CDCl₃) δ 2.31(d, 1H), 2.4(dd, 2H), 2.75(dd, 2H),
 2.85(dd, 1H), 3.36(br. s, 1H), 3.45(br. s, 1H), 3.6(br. t, 2H), 3.82(br. m, 2H), 3.95(br. d, 2H), 4.35(m, 2H),
 4.42(d, 1H), 4.55(m, 1H), 4.70(d, 1H), 4.82(br. s, 2H),
 10 5.5(d, 1H), 6.91(d, 1H), 7.25(br. m, 5H), 7.35-7.46(br.
 - (3S)-3-[(3S)-2-0xo-3-benzoylformylamino-5-(2-hydroxy)acetyl-2,3,4,5-tetrahydro-1H-1,5-

m, 3H), 7.5-7.6(m, 2H), 8.15(br. d, 2H).

benzodiazepine-1-acetylamino]4-oxo-butyric acid (681),

15 was synthesized from **680** by the method used to prepare **678** from **677** to afford 45 mg of **681** as a grey solid, 1 H NMR (CD₃OD) δ 2.5(m, 1H), 2.7(dt, 1H), 3.55-3.85(br. m, 3H), 4.05(m, 1H), 4.3(m, 1H), 4.5-4.7(br. m, 3H),

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4.85(br. s, 2H), 7.3(br. m, 2H), 7.4-7.7(m, 5H), 8.15(d, 2H).

(3s)-2-Oxo-3-benzoylamino-5-(2-acetoxy)acetyl-N[(2Rs,3s)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]5 2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide
(682), was synthesized from 600b by the methods used to
prepare 655 from 600b to afford 495 mg of 682 as a
white solid, ¹H NMR (CDCl₃) & 2.00(s, 3H), 2.05(s, 3H),
2.47(d, 1H), 2.58(dd, 1H), 2.85(dd, 1H), 2.89(dd, 1H),
10 3.9(m, 2H), 4.05-4.15(m, 2H), 4.19(dd, 1H), 4.45(m,
2H), 4.55-5.05(m, 8H), 5.55(d, 1H), 6.85(d, 1H),

(3S)-3-[(3S)-2-Oxo-3-benzoylamino-5-(2-acetoxy)acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-

7.15(d, 1H), 7.25-7.55(m, 10H), 7.75(d, 2H).

- 20 4.6(d, 1H), 4.72(d, 1H), 4.95(br. s, 2H), 7.45(zr. m, 2H), 7.52-7.65(br. m, 5H), 7.88(d, 2H).

10 (3S)-2-0xo-3-(3-chloro-4-aminobenzoyl) amino-5-(2triisopropylsilyloxy) acetyl-N-[(2RS,3S)-benzyloxy-5oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-

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benzodiazepine-1-acetamide (685), was synthesized from 600b by the methods used to prepare 676 from 600b to afford 165 mg of 685.

(3S)-3-[(3S)-2-0xo-3-(3-chloro-4-aminobenzoyl) amino-55 (2-triisopropylsilyloxy) acetyl-2,3,4,5-tetrahydro-1H1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid
(686). To a solution of 685 (165 mg, 0.21 mmol) in THF
was added a solution of TBAF (1M, 0.21 mL). The
product was isolated by filtration after precipitation
10 from reaction mixture. Reverse phase corromatography
(10% to 80% MeCN in water/ 0.1% TFA) afforded 25 mg of
686 as a white solid, ¹H NMR (CD₃OD) δ 2.37-2.42 (m),
2.59-2.70 (m), 3.60-3.89 (m), 4.01 (d), 4.20-4.31 (m),
4.42-4.70 (m), 4.80-5.05 (m), 6.79 (d), 7.32-7.65 (m),
15 7.81 (s).

(3S)-2-0xo-3-(3,5-dichloro-4-hydroxybenzoyl)amino-5-methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid (687a), was synthesized from 600b using methods similar to those used for preparing 654 from 5 600b to afford 1.6 g of 687a.

- (3S)-2-0xo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetic acid (687b), was synthesized from 600b using methods similar to those used for preparing 654 from
- 10 600b to afford 1.1 g of 687b.
 - (38) -2-0xo-3-(3,5-dichloro-4-hydroxybenzoyl)amino-5-methoxyacetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (688a). To a solution of
- 15 (3S,2R,S)-3-allyloxycarbonylamino-2-benzyloxy-5oxotetrahydrofuran (Chapman, Biorg. Med. Chem. Lett.,
 2, pp. 613-618 (1992)) (1.13 g, 1.2 equiv) in CH₂Cl₂
 was added triphenylphosphine (423 mg, 0.5 equiv),
 dimethylbarbituric acid (1.26 g, 2.5 equiv), and
- 20 tetrakistriphenylphosphine palladium (0) (373 mg, 0.1 equiv). After 5 minutes the reaction mixture was cooled via ice-bath then added a solution of 687a in DMF (1.6 g, 1 equiv), HOBT (460 mg, 1.1 equiv), and EDC (681 mg, 1.1 equiv). The resulting mixture was allowed.
- 25 to stir at ambient temperature. After 16 hours the reaction mixture was poured into NaHSO₄ and extracted twice with EtOAc. The organic layer was washed with NaHCO₃, brine, dried over Na₂SO₄ and concentrated in vacuo. Chromatography (SiO₂, 20% to 100% EtOAc in
- 30 CH2Cl2) afforded 880mg of 688a as an off-white solid, "H

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NMR (CD₃OD) δ 2.55(dd, 1H), 2.7(dd, 1H), 3.0(m, 1H), 3.6(m, 1H), 3.75(d, 1H), 3.9-4.0(m, 2H), 4.3-4.45(m, 3H), 4.5-4.6(m, 3H), 4.7(m, 2H), 5.35(s, 1H), 5.55(d, 1H), 7.1-7.5(m, 4H), 7.85(s, 2H).

- 5 (3S)-2-Oxo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-methoxyacetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (688b), was synthesized from 687b by the method used to prepare 688a from 687a to afford 960 mg of 688b as an off-white solid, ¹H MMR (CD₃OD) δ 2.6(dd, 1H), 2.7(dd, 1H), 3.0 (dd, 1H), 3.2(s, 3H), 3.7 (m, 3H), 3.9 (m, 2H), 4.4-4.5 (m, 2H), 4.6 (m, 3H), 5.35(s, 1H), 5.55(d, 1H), 7.25 (m, 2H), 7.4-7.5 (m, 4H).
- 15 (3S)-3-[(3S)-2-0xo-3-(3,5-dichloro-4-hydroxybenzoyl)amino-5-methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (689a), was synthesized from 688a by the method used to prepare 2002 from 2001 to afford 184 mg
- 20 of 689a as a white solid, 1 H NMR (CD₃OD) δ 2.45(m, 1H), 2.6(m 1H), 3.3(s, 3H), 3.7-3.85(m, 2H), 4.0(d, 1H), 4.3(m, 1H), 4.5-4.6(m, 3H), 7.3-7.6(m, 4H), 7.85(s, 2H).

(3S) -3 - [(3S) -2 -0xo -3 - (3,5 -dimethyl -4 -

25 hydroxybenzoyl)amino-5-methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (689b), was synthesized from 688b by the method used to prepare 2002 from 2001 to afford 412 mg of 689b as a white solid, ¹H NMR (CD₃OD) δ 2.5(m, 1H).

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2.7(m, 1H), 3.3(s, 3H), 3.7-3.85(m, 2H), 4.05(dd, 1H), 4.3(m, 1H), 4.6(m, 2H), 7.45-7.4(m, 2H), 7.5(s, 2H), 7.55(m, 2H).

(3S)-2-Oxo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5hydroxyacetyl-N-[(2RS,3S)-benzyloxy-5-oxotetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5benzodiazepine-1-acetamide (690a), was synthesized from
600b via methods used to prepare 676 from 600b, 688a
from 687a, then 677 from 676 to afford 863 mg of 690a

10 as a white solid, ¹H NMR (CD₃OD) δ 2.2(s, 6H), 2.45(d,
0.5H), 2.6-2.9(m, 1H), 3.05(dd, 0.5H), 3.65-3.85(m,
2H), 3.95-4.1(m, 1H), 4.35-5.0(m, 7H), 5.35(s, C.5H),

(3S)-2-0xo-3-(4-hydroxybenzoyl)amino-5-hydroxyacetyl-N[(2Rs,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide
(690b), was synthesized from 600b via methods used to
prepare 677 from 600b to afford 200 mg of 690b. 1H NMR

5.65(d, 0.5H), 7.2-7.4(m, 4H), 7.4-7.7(m, 7H).

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 $\begin{array}{c} (\text{CD}_3\text{OD}) \;\; \delta \;\; 2.49\,(\text{d},\;\; 1\text{H}),\;\; 2.65\,(\text{d},\;\; 1\text{H}),\;\; 2.86\,(\text{d},\;\; 1\text{H}),\;\; 2.85\,(\text{d},\;\; 1\text{H}),\;\; 2.87\,(\text{d},\;\; 1\text{H}),\;\; 3.05\,(\text{dd},\;\; 1\text{H}),\;\; 3.35\,(\text{br. s},\;\; 1\text{H}),\;\; 3.72\,(\text{br. s},\;\; 2\text{H}),\;\; 4.01\,(\text{m},\;\; 2\text{H}),\;\; 4.45\,(\text{br. m},\;\; 1\text{H}),\;\; 4.6\,(\text{m},\;\; 1\text{H}),\;\; 4.7\,(\text{m},\;\; 1\text{H}),\;\; 4.8\,(\text{m},\;\; 1\text{H}),\;\; 4.95\,(\text{br. s},\;\; 2\text{H}),\;\; 5.65\,(\text{d},\;\; 1\text{H}),\;\; 6.8\,(\text{d},\;\; 2\text{H}),\;\; 7.2-7.35\,(\text{br. m},\;\; 3\text{H}),\;\; 7.45\,(\text{m},\;\; 2\text{H}),\;\; 7.75\,(\text{d},\;\; 2\text{H}),\;\; \end{array}$

(3S)-3-[(3S)-2-Oxo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-hydroxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (691a), was synthesized from 690a by the method used to prepare 2002 from 2001 to afford 560 mg of 691a as a white solid, ¹H NMR (CD₃OD) δ 2.15(s, 6H), 2.45(m, 1H), 2.65(m, 1H), 3.55(m, 1H), 3.7(d, 1H), 4.0(d, 1H), 4.25(m, 1H), 4.5-4.6(m, 3H), 7.3-7.5(m, 15 6H).

(3S)-3-[(3S)-2-0xo-3-(4-hydroxybenzoy1)amino-5-hydroxyacety1-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (691b), was synthesized from 690b by the method used to prepare 20 2002 from 2001 to afford 410 mg of 691b as a white solid, ¹H NMR (CD₃OD) δ 2.5 (m, 1H), 2.65 (m, 1H), 3.75 (m, 1H), 3.8 (d, 1H), 4.05 (d, 1H), 4.25 (m, 1H), 4.5 (m, 1H), 4.6 (m, 1H), 4.95 (br. s, 2H), 6.8 (d, 2H), 7.45 (m, 2H), 7.6 (m, 2H), 7.75 (d, 2H).

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(3S)-2-Oxo-3-benzoylamino-5-hydroxyacetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (695a), was synthesized from 600b via methods used to prepare 5 677 from 600b to afford 75 mg of 695a, ¹H NMR (CD₃OD) δ 2.2(s, 6H), 2.45(m, 1H), 2.6(m, 1H), 3.65(m, 1H), 3.75(d, 1H), 4.0(d, 1H), 4.28(m, 1H), 4.5(m, 3H), 7.4-7.6(m, 6H).

(3S) -2-Oxo-3-(4-acetamidobenzoyl)amino-5-hydroxyacetyl10 N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide
(695b), was synthesized from 600b via methods used to
prepare 677 from 600b to afford 880 mg of 695b, ¹H NMR
(CDCl₃) δ 2.1(s, 3H¹, 2.25-2.5(m, 2H), 2.8-2.92(m,

15 0.5H), 3.15-3.2(m, 0.5H), 3.45-3.6(m, 2H), 3.75-3.95(m, 2H), 4.15-4.25(m, 1H), 4.35-4.6(m, 2H), 4.6-4.88(m, 3H), 5.22(s, 0.25H), 5.33(s, 0.25H), 5.52-5.58(d, 0.5H), 7.15-7.45(m, 9.5H), 7.5-7.75(m, 5H), 8.3-8.35(m, 0.5H), 9.08-9.18(m, 1H).

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(3S) -2RS-Oxo-3-(3,5-dimethyl-4-hydroxybenzoyl) amino-5-hydroxyacetyl-N-(2-benzyloxy-5-oxo-tetrahydrofuran-3-yl)-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (695c), was synthesized from 600b via methods used to prepare 677 from 600b to afford 840 mg of 695c, ¹H NMR(CDCl₃) δ 2.23(s, 3H), 2.26(s, 3H), 2.45-2.62(m, 1H), 2.8-2.9(dd, 0.5H), 2.9-3.05(dd, 0.5H), 3.45-3.63(m, 1H), 3.64(s, 1.5H), 3.68(s, 1.5H), 3.78-4.05(m, 2H), 4.2-4.33(m, 1H), 4.4-4.63(m, 2H), 4.65-4.94(m, 2H), 4.95-5.1(m, 1H), 5.45(s, 0.5H), 5.5-5.6(dd, 0.5H),

6.9-6.95(d, 1H), 7.25-7.7(m, 12H).

(3S)-2-0xo-3-(3,5-dichloro4-hydroxybenzoy1)amino-5hydroxyacety1-N-[(2RS,3S)-benzyloxy-5-oxotetrahydrofuran-3-y1]-2,3,4,5-tetrahydro-1H-1,5-

- benzodiazepine-1-acetamide (692a), was synthesized from 600b via methods used to prepare 661 from 600b, excluding steps used to make 604d from 603d, using instead the method to prepare 688a from 687a to afford 854 mg of 692a, ¹H NMR (CD₃OD) & 2.45(d, 1H), 2.6(m,
- 20 1H), 2.7(m, 1H), 3.0(m, 1H), 3.5-3.7(m, 4H), 4.6(q, 2H), 4.45(m, 3H), 4.55(m, 4H), 5.35(s, 1H), 5.6(d, 1H), 7.2-7.5(m, 9H), 7.85(s, 2H).

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(3S)-2-0xo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-hydroxyacetyl-N-[(2RS,3S)-ethoxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (692b), was synthesized from 600b via methods used to prepare 661 from 600b, excluding steps used to make 604d from 603d, using instead the method to prepare 688a from 687a to afford 207 mg of 692b, ¹H NMR (CD₃OD) δ 1.05(t, 3H), 1.15(t, 3H), 2.45(d, 1H), 2.55(m, 1H), 2.7 (m, 1H), 3.55(m, 2H), 3.6-3.75(m, 5H), 10 4.0(dd, 2H), 4.3(d, 1H), 4.4-4.7 (m, 5H), 5.25(s, 1H), 5.5(d, 1H), 7.25-7.6 (m, 4H), 7.85(s, 2H).

(3s)-2-Oxo-3-benzoylamino-5-acetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (693), was synthesized from 600b via methods used to prepare 688a from 600b to afford 30 mg of 693, ¹H NMR (CD₃OD) δ 1.7(s, 3H), 1.8(s, 3H), 2.51(d, 1H), 2.6(m, 1H), 2.85(m, 1H), 3.0(m, 1H), 3.75(br. d, 2H), 4.0-4.1(dd, 2H), 4.5-5.0(m, 6H), 5.45(s, 1H), 5.55(s, 1H), 7.15-20 7.85(m, 14H).

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(35) -3-[(35) -2-0xo-3-(3,5-dimethyl-4-methoxybenzoyl)amino-5-hydroxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (694), was synthesized from 691c by the 5 method used to prepare 2002 from 2001 to afford 380 mg of 694 as a white solid, ¹H NMR (CD₃OD) & 2.25(s, 6H), 2.45(m, 1H), 2.65(m, 1H), 3.65(m, 5H), 4.0(d, 1H), 4.28(m, 1H), 4.55(d, 2H), 4.95(m, 1H), 7.4-7.6(m, 6H).

Compounds 700-711 were prepared by methods

10 similar to the methods used to prepare compounds 619635 (see, Example 13). Physical data for compounds
700-711 is listed in Table 25.

Compounds 910-915 and 918-921 were prepared as described below. Physical data for these compounds is 15 listed in Table 26.

Structure

MS (M+Na)+	575.9	572.1
HPLC RT min (method)	15.855 (1)	10.315 (2)
MM	552.50	547.53
MF	C26H24N4O10	C27H25N5O8
Structure		
Compound	702	703

	T		
MS (M+Na)+	562.1	562.1	592.4
HPLC RT min (method)	10.475 (2)	14.260 (1)	14.836 (1) 97%
ММ	538.52	538.52	568.55
MF	C2 6H2 6N409	C2 6H2 6N409	C27H28N4010
Structure			
Compound	704	705	706

				HPLC RT min	
Compound	Structure	MF	MW	(method)	MS
				Purity	+ (B+Na)
707	CONTRACTOR TO THE CONTRACTOR T	C27H28N409	552.55	15.952 (1) 988	575.9
708		C27H26N409	550.53	10.731 (2) 93%	574.6
709		C28II30N408	550.57	13.192 (2) 953	574

T	T
582.2	521.9
12.406 (2)	13.072 (1)
557.95	498.45
C25H24C1N508	C23H22N409
710	711
	CZSHZ4CIN508 557.95 12.406 (2)

-	716	

MS (M+Na)+	564.4	5.77.5
HPLC RT min (method)	8.172 (2) 99%	6.949 (2) 99%
MM	540.49	553.53
MF	C25H24N4O10	C2 6H2 7N509
Structure	DE LOS DEL LOS DELLOS DE LOS DELLOS DE LOS DELLOS D	
Compound	910	911

Table 26

Compound	0 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	174		HPLC RT min	MS
	a range	Ξ	M M	(method)	(M+Na) +
912	OH OF THE CONTRACT OF THE CONT	C25H26N409	526.51	8.317 (2) 99%	550.7
o 11 60	O Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	C26H29N5OB	539.55	6.588 (2)	563.5

				HPLC RT min	044
Compound	Structure	MF	MM	(method)	MS + (M+Na) +
				Purity	
914		C26H26C1N509	587.98	7.815 (2) 99%	612.2
915		C26H25C12N5O9 622.42	622.42	7.490 (2)	647

				HPLC RT min	
Compound	Structure	MF	MM	(method)	MS
				Purity	(M+Na)+
916/691b	HO N HO HO HO	C24H24N409	512.48	6.331 (2) 98%	537
917/691a	OF THE STATE OF TH	C26H28N409	540.53	8.114 (2)	564.9

		T	
MS (M+Na)+	619.3		
HPLC RT min (method) Purity	11.817 (2)	9.709 (2) 91%	
ММ	595.40	535.52	
ΜF	CZ5HZ4Cl2N409 595.40		
Structure	H ₂ C-O _C H		
Compound	918	919	

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MS (M+Na)+	560.6	579.1
HPLC RT min (method)	5.494 (2)	7.827 (2)
MM	536.51	554.52
MF	C25H24N6O8	C26H26N4O10
Structure		OF TO STATE OF THE
Compound	920	921

				HPLC RT min	-
	Structure	MF	ΜW	(method)	M. M.
				Purity	+ (M+M)
- i	H ₂ C ₂ C ₃ H	C27H30N409	554.56	554.56 10.024 (2)	578.8

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Step A. Synthesis of 401. TentaGel S® NH2 resin (0.25 mmol/g, 6.8 g) was placed in a glass shaker vessel and washed with dimethylacetamide (3 X 20 mL). To a solution of 400 (1.70 g, 2.9 mmol, prepared from 5 (3S) 3-(fluorenylmethyloxycarbonyl)-4-oxobutryic acid t-butyl ester according to A.M. Murphy et. al. J. Am. Chem. Soc., 114, 3156-3157 (1992)) in dimethylacetamide (15 mL) was added O-benzotriazole-N,N,N,N'tetramethyluronium hexafluorophosphate (HBTU: 1.09 g. 10 2.9 mmol), and DIEA (1.0 mL, 5.7 mmol). The solution was added to the resin, followed by dimethylacetamide (5 mL). The reaction mixture was agitated for 3 h at room temperature using a wrist arm shaker. The resin was isolated by suction filtration and washed with 15 dimethylacetamide (6 X 20 mL). A sample of resin (7.4 mg) was thoroughly washed with 50% methanol in dichloromethane and dried under suction. Deprotection of the Fmoc group using 20% piperidine in dimethylacetamide (10.0 mL) and UV analysis of the 20 solution revealed a substitution of 0.19 mmol g-1.

Step B. Synthesis of 903. Resin 401 was deprotected with 20% (v/v) piperidine/dimethylacetamide (20 mL) for 10 min (shaking) and then for 10 min with fresh piperidine reagent (20 ml). The resin was then washed with dimethylacetamide (6 X 20 ml). A solution of 902 (1.52 g, 2.81 mmol) was treated with HBTU (1.07 g, 2.83 mmol) and DIEA (1.0 mL, 5.7 mmol) and transferred to the resin, followed by dimethylacetamice (5 mL). The reaction mixture was agitated for 2.5 h at room temperature using a wrist arm shaker. The resin was isolated by suction filtration and washed with

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dimethylacetamide (4 X 20 mL) and dichloromethane (4 X 20 mL), and dried under nitrogen purge. Resin substitution was performed as described for 401 and determined to be 0.169 mmol $\rm g^{-1}$.

5

Step C. Synthesis of 905. Resin 903 (7.54 q, 1.27 mmol) and dimedone (2.19 g, 15.6 mmol) were placed in a 100 mL round bottomed flask and freshly distilled anhydrous tetrahydrofuran (60 mL) was added.

10 Tetrakis(triphenylphosphine)palladium (0) (0.32 g, 0.28 mmol) was added and the nitrogen blanketed, sealed reaction was agitated for 15 h on a wrist action shaker. The resin was filtered, washed with dimethylacetamide (4 X 20 mL), dichloromethane (4 X 20 mL) and dimethylacetamide (1 X 20 mL). Sufficient dimethylacetamide was added to the resin to obtain a slurry followed by pyridine (1.5 mL, 18.5 mmol) and a solution of 904 (5.5 mmol) in dichloromethane (10 mL).

The reaction was shaken under nitrogen for 8 h, then 20 filtered. The resin was washed with dimethylacetamide $(5 \times 20 \text{ mL})$ and dichloromethane $(5 \times 20 \text{ mL})$.

Step D. Synthesis of 906. This compound was prepared from resin 905 (0.24 g, 0.038 mmol) using an 25 Advanced ChemTech 396 Multiple Peptide synthesizer. The automated cycles consisted of a resin wash with dimethylformamide (3 X 1 mL), deprotection with 25° (v/v) piperidine in dimethylformamide (1 mL) for 10 min followed by fresh reagent (1 mL) for 20 min to yield resin 906. The resin was washed with dimethylformamide (3 X 1 mL) and N-methypyrrolidone (3 X 1 mL).

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Step E. (910-922) Resin 906 was acvlated with a solution of 0.4M carboxylic acid and 0.4M HOBT in Nmethypyrrolidone (0.5 mL), a solution of 0.4M HBTU in N-methylpyrrolidone (0.5 mL) and a solution of 1.6M 5 DIEA in N-methypyrrolidone (0.25 mL) and the reaction was shaken for 2 hr at room temperature. The resin was washed with N-methylpyrrolidone (1 X 1 mL), dimethylformamide (4 X 1 mL), 50% methanol in dichloromethane (5 X 1 mL) and dried in air. The 10 aldehyde was cleaved from the resin and globally deprotected by treatment with 95% TFA/ 5% H2O (v/v, 1.5 mL) for 30 min at room temperature. After washing the resin with cleavage reagent (2 X 1 mL), the combined filtrates were added to cold 1:1 ether:hexane (35 mL) 15 and the resulting precipitate was isolated by centrifugation and decantation. The resulting pellet was dissolved in acetonitrile (0.5 mL) and H2O (0.5 mL) and filtered through 0.45 micron microcentrifuge

20 RP-HPLC with a Rainin Microsorb^M C18 column (5 μ, 21.4 X 250 mm) eluting with a linear acetonitrile gradient (10% - 50%) containing 0.1% TFA (v/v) over 30 min at 12 mL/min. Fractions containing the desired product were

filters. The compound was purified by semi-preparative

pooled and lyophilized to provide 910-922.

Analytical HPLC methods:

25

(1) Waters DeltaPak C18, 300Å (5µ, 3.9 X 150 mm).
Linear acetonitrile gradient (0% - 25%) containing 0.1
TFA (v/v) over 14 min at 1 mL/min.

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(2) Waters DeltaPak C18, 300Å (5 μ , 3.9 X 150 mm). Linear acetonitrile gradient (5% - 45%) containing 0.1% TFA (v/v) over 14 min at 1 mL/min.

(3S) -3-[(3S) -2-0xo-3-(isoquinolin-1-oyl)amino-5-

5 hydroxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxo-butyric acid (696) was synthesized from 600b by the method used to prepare 691a from 600b to afford 696.

1H NMR (CD3OD) δ 2.45(m, 1H), 2.7(m, 1H), 3.75(d, 1H), 3.95(q, 1H), 4.05(d, 1H), 4.3(m, 1H), 10 4.45-4.65(m, 2H), 5.05(m, 1H), 7.5-7.6(m, 3H), 7.7(t, 1H), 7.8(t, 1H), 7.98(t, 1H), 8.55(d, 1H), 9.1(d, 1H).

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(3S)-2-Oxo-3-(isoquinolin-1-oyl)amino-5-hydroxyacetyl-N-[(2RS,3S)-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (696a) was synthesized from 600b via methods used to prepare 690a from 600b to afford 696a. ¹H NMR (CDCl₃) δ 0.95(t, 2H), 1.25(t, 1H), 1.4(m, 2H), 1.55(m, 1H), 2.55(m, 1H), 2.85(m, 1H), 2.95(dd, 1H), 3.15(m, 1H), 3.55(m, 1H), 3.9(m, 2H), 4.35(t, 1H), 4.4-4.55(m, 2H), 4.75(m, 1H), 4.8-5.05(m, 2H), 5.55(d, 1H), 5.55(d, 1H), 7.15(d, 1H), 7.2-7.5(m, 5H), 7.6-7.8(m, 3H), 8.45(d, 1H), 9.05(d, 1H), 9.35(d, 1H).

(3S)-2-Oxo-3-(isoquinolin-1-oyl)amino-5-hydroxyacetyl-N-[(2RS,3S)-ethoxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-carboxamide (696b)

15 was synthesized from 600b via methods used to prepare 690a from 600b to afford 696b. ¹H NMR (CDCl₃) δ 0.9(m, 3H), 1.15(q, 3H), 1.15(m, 1H), 1.65(m, 1H), 2.5(m, 1H), 2.8(m, 1H), 2.95-3.0(m, 2H), 3.6(m, 2H), 3.7-3.85(m, 4H), 4.0(m, 2H), 4.3(m, 1H), 4.55(m, 1H), 5.45(d, 1H), 4.65-4.95(m, 1H), 5.05(m, 1H), 5.35(s, 1H), 5.45(d, 1H), 6.85(d, 1H), 7.25(d, 1H), 7.35-7.85(6H), 8.85(dd, 2H), 9.05(m, 1H), 9.35(dd, 2H).

[2RS-(4-chlorobenzyl)oxy-5-oxo-tetrahydrofuran-3-yl]25 2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-carboxamide
(696c) was synthesized from 600b via methods used to
prepare 690a from 600b to afford 696c. ¹H NMR (CD30D) δ

1.25(t, 1H), 1.65(q, 1H), 1.9(m, 1H), 2.9(m, 1H),

(3S) ~2-0xo-3-(isoquinolin-1-oyl)amino-5-hydroxyacetyl-

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3.05(m, 1H), 3.9(d, 1H), 4.2(m, 1H), 4.3(d, 1H), 4.7-5.0(m, 3H), 5.25(m, 1H), 5.7(s, 1H), 5.9(d, 1H), 7.5(d, 2H), 7.7-7.9(m, 3H), 8.0(t, 1H), 8.2(m, 2H), 8.75(d, 1H), 9.35(d, 1H).

- 5 (3S)-2-Oxo-3-(isoquinolin-1-oyl)amino-5-hydroxyacetyl(2RS-cyclopentyloxy-5-oxo-tetrahydrofuran-3-yl)2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-carboxamide
 (696d) was synthesized from 600b via methods used to
 prepare 690a from 600b to afford 696d.

 1 h NMR (CDCl₃) δ

 10 0,9(t, 1H), 1.2(t, 1H), 1.3-1.45(m, 2H), 1.6-1.8(m,
 4H), 2.45(m, 1H), 2.8(m, 1H), 3.0(m, 1H), 3.4(q, 1H),
 3.5(d, 1H), 4.0(m, 2H), 4.2-4.3(m, 2H), 4.55(d, 1H),
 4.65(m, 1H), 4.9(m, 1H), 5.05(m, 1H), 5.4(s, 1H),
 5.5(d, 1H), 6.8(d, 1H), 7.3-7.9(m, 6H), 8.5(d, 1H),
 15 9.05(d, 1H), 9,4(d, 1H).
 - (3S)-2-0xo-3-(isoquinolin-1-oyl)amino-5-hydroxyacetyl-N-[(2R,3S)-phenethoxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide (696e) was synthesized from 600b via methods used to
- 20 prepare 690a from 600b to afford 696e. ¹H NMR (CDCl₃) 8 1.2(t, 1H), 2.4(m, 1H), 2.8(m, 2H), 3.6(d, 1H), 3.7(q, 1H), 4.0(m, 2H), 4.3(d, 2H), 4.65(m, 1H), 4.85(t, 1H), 5.0(m, 1H), 5.35(d, 1H), 6.5(d, 1H), 7.15-7.85(m, 8H), 8.45(d, 1H), 9.05(d, 1H), 9.4(d, 1H).

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Example 32

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20

Compound	UV- Visible Ki (nM)	Cell PBMC avg. IC50 (nM)	Whole human blood IC50 (nM)	Clearance Mouse, i.v. ml/min/kg	Clearance Rat, i.v. ml/min/kg
688c	200				
689b-1	3.5		2700		
696-1	0.5				
696-2	0.5				
697	1.8		5000		
698	18		13500		
699	1.1				
699a-2					
720	2.7				
721	1.3		5000		
722	5		5000		
723	2.3		2000		
724	2		1800		
725	3.7		3000		
726	300				
727	50		2300		
728	300				
729	28		2800		
730	90		8000		
731	150				
732	5		1800		
733	5		1500		
734	9		6000		
735	6		10000		

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Example 33

Compounds 684a, 688b-1, 688c, 689b-1, 690a-1, 696-1, 696-2, 696a-2, 696a-1, 697a, 698a, 698a, 699, 699a, 699a-1, 699a-2, 800 and 801 were prepared as 5 described below.

Table 28

CIP#	R ⁴	R ³	R ⁵	R ¹
684a	E CHO	CH3	Н	OtBu
688b-1	CH ₃	MeO	F	OBn
688c	CH ₃	MeO	Н	
689b-1	CH ₃	MeO_j	F	OH H
690a-1	HO CH	HQ HQ	Н	OEt OEt

CIP#	R ⁴	R ³	R ⁵	R ¹
696-1	Ë	но	F	O H
696-2	r. J.	ющ	Cl	0=(= 0
696a-2	90	ф Но	Cl	OOBn
696a-1	£	PO T	F	O O O O O O O O O O O O O O O O O O O
697	H ₂ N G	9 9	Н	0=(_0 _E 0
697a	CI NO	H	Н	OBn
698		± >=0	Н	of p
698a	j	± >=0	Н	OBn
699	(F).	MeO (Н	OH H
699a	F.	№	Н	OBn

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CIP#	R ⁴	R ³	R ⁵	R ¹
699a-1	° JO	MeO_ii	F	OBn
699a-2	i di	MeO	F	O H O H
800	YS OF	H0 =	Н	OOBn
801	S. S) S	Н	0 H

5 (3S)-3-[(3S)-2-0xo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-hydroxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4,4-diethoxybutyric acid ethyl ester(690a-1), was synthesized by the methods used to prepare 690a and 10 2100b to afford 690a-1, \(^1\)H NMR(CDCl₃) \(^5\) 1.15(t, 6H), 1.3(t, 3H), 2.25(s, 6H), 2.60(d, 2H), 3.50(m, 2H), 3.70(m, 4H), 4.05(m, 2H), 4.15(m, 2H), 4.30(d, 1H), 4.45(m, 1H), 4.50(d, 1H), 4.55(d, 1H), 4.70(t, 1H), 5.05(m, 1H), 5.30(s, 1H), 6.70(d, 1H), 7.10(d, 2H), 15 7.30-7.50(m, 7H)

(3S)-2-Oxo-3-(3,5-dichloro-4-aminobenzoyl)amino-5-hydroxyacetyl-N-[(2RS,3S) 2-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetamide(697a) was synthesized via 20 methods used to prepare 677 to afford 840 mg of 697a,

1H NMR (CDCl₃) 8 1.78 (br. s, 2H), 2.48-2.58 (d, 0.5H),

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2.6-2.7 (m, 0.5H), 2.8-2.9 (m, 0.5H), 2.92-3.03 (m, 0.5H), 3.55-3.8 (m, 2H), 3.92-4.02 (d, 1H), 4.25-4.3 (d, 0.5H), 4.37-4.42 (d, 0.5H), 4.43-4.48 (m, 0.5H), 4.55-4.65 (m, 1.5H) 4.7-5.12 (m, 5H), 5.44 (s, 0.5H), 5.58-5.63 (d, 0.5H), 6.95-8.1 (m, 13H).

(3S)-3-[(3S)-2-0xo-3-(3,5-dichloro4-aminobenzoy1)amino-5-acetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxobutyric acid (697) was synthesized via methods used to prepare 2002 from 2001 to afford 140 mg 10 of 697, ¹H NMR (CD₃OD) & 238-2.5 (m,1H), 2.55-2.75 (m, 1H), 3.68-3.9 (m, 3H), 3.95-4.03 (m, 1H), 4.2-4.3 (m, 1H), 4.4-4.7 (m, 4H), 7.35-7.8 (m, 6H).

(3S)-3-[(3S)-2-Oxo-3-(3,5-dimethyl-4-methoxybenzoyl)amino-5-hydroxyacetyl-2,3,4,5
15 tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-acetoxy-3-butenoic acid ethyl ester(684a), was synthesized by the methods used to prepare 2100j to afford 684a, ¹H NMR (500 MHz, CDCl₃ mixture of diastereomers) δ 1.3 (s, 9H), 1.6(s, 3H), 2.1(s, 3H), 2.1(s, 3H), 2.1(s, 3H), 3.9(m, 1H), 4.1(d, 1H), 4.3(d, 1H), 4.6-4.8(m, 3H), 3.9(m, 1H), 6.7(s, 1H), 7.0(d, 1H), 7.1(d, 1H), 7.2-7.5(m, 6H).

(3S) -2-0xo-3-isoquinolin-1-oylamino-5-formyl-N-

25 [(2Rs,3s) 2-benzyloxy-5-oxo-tetrahydrofuran-3-yl]2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1acetamide(698a) was synthesized via methods used to
prepare 652 to afford 795 mg of 698a ¹H NMR (500 MHz,
CDCl₃ mixture of diastereomers) δ 2.8(m, 2H), 4.0(m,
30 IH:, 4.5-4.8(m, 4H), 5.2(m, 1H), 5.5(s, 1H), 5.75(d,

- 735 -

1H), 7.3-7.85 (m, 11H), 7.9 (t, 1H), 8.2 (d, 1H), 8.6 (m, 1H), 9.3 (m, 1H).

- (3S)-3-[(3S)-2-0xo-3-isoquinolin-1-oylamino-5-formyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-
- 5 acetylamino]4-oxobutyric acid(698) was synthesized via methods used to prepare 653 to afford 225 mg of 698 $^{1}\mathrm{H}$ NMR (500 MHz, CD₃OD) δ 2.4(m, 1H), 2.6(m, 1H), 3.9(m, 1H), 4.2(m, 1H), 4.3-4.7(m, 4H), 5.1(m, 1H), 7.3-7.5(m, 4H), 7.6-7.8(m, 2H), 7.8(m, 2H), 8.2(d, 1H), 8.5(d,
- 10 lH), 9.0(d, lH).
 - (3S)-2-0xo-3-isoquinolin-1-oylamino-5-methoxyacetyl-N-[(2RS,3S) 2-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-
- acetamide(699a) was synthesized via methods used to
 15 prepare 655 to afford 820 mg of 699a as a tan solid, ¹H
 MMR (500 MHz, CDCl₃) 5 2.60 (ddd, 1H), 2.90 (ddd, 1H),
 3.20 (s, 3H), 3.25 (s, 3H), 3.70 (t, 1H), 3.90 (m, 2H),
 4.20 (dd, 1H), 4.60 (m, 2H), 4.70-5.00 (m, 5H), 5.55
 (d, 1H), 7.00 (d, 1H), 7.20-7.50 (m, 7H), 8.45 (dd,
- 20 1H), 9.0 (dd, 1H), and 9.35 ppm (dd, 1H).
 - (3S)-2-0xo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-methoxyacetyl-N-[(2RS,3S) 2-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-7-fluoro-1H-1,5-benzodiazepine-1-acetamide(688b-1) was synthesized via methods used to prepare 655 to afford 600 mg of
- 25 via methods used to prepare 655 to afford 600 mg of 688b-1, $^{1}{\rm H}$ MMR (CDCl $_{3}$; mix. of diastereomers) δ 2.21 (s, 3H), 2.28 (s, 3H), 2.42-2.50 (m, 0.5 H), 2.58-2.65 (m, 0.5H), 2.83-2.91 (m, 0.5H), 2.98-3.1 (m, 0.5H), 3.18 (s,1.5H), 3.22 (s, 1.5H), 3.72-3.78 (d, 1H), 3.78-
- 30 3.9 (m, 2H), 4.08-4.15 (d, 1H), 4.5-4.69 (m, 3H), 4.7-

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5 (3s)-3-{(3s)-2-Oxo-3-(3,5-dimethyl-4-hydroxybenzoyl)amino-5-methoxyacetyl-2,3,4,5-tetrahydro-7-fluoro-1H-1,5-benzodiazepine-1-acetylamino|4-oxobutyric acid(689b-1) was synthesized via methods used to prepare 2002 from 2001 to afford 10 689b-1, ¹H NMR (CD₃OD) & 2.18 (s, 6H), 2.36-2.47 (m, 1H), 2.6-2.72 (m, 1H), 3.34 (s, 3H), 3.66-3.88 (m, 2H), 3.95-4.05 (m, 1H), 4.2-4.78 (m, 5H), 4.9 (m, 1H), 7.3-7.41 (m, 2H), 7.48 (s, 2H), 7.5-7.63 (m, 1H).

(3S) -3-[(3S)-2-0xo-3-isoquinolin-1-oylamino-5-

- 15 methoxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine1-acetylamino]4-oxobutyric acid(699) was synthesized via methods used to prepare 2002 from 2001 to afford 699 as a white solid, ¹H NMR (500 MHz, CD₃OD) δ 2.50 (m, 1H), 2.70 (m, 1H), 3.25 (s, 3H), 3.80 (bd, 1H), 20 3.90 (bd, 1H), 4.00 (bd, 1H), 4.30 (m, 1H), 4.50-4.70 (m, 3H), 4.80-4.85 (bt, 1H), 5.00 (bm, 1H), 7.40-7.55 (m, 5H), 7.70 (bm, 1H), 7.85 (bm, 1H), 8.00 (bm, 1H),
 - 8.55 (bd, 1H), and 9.05 ppm (bd, 1H).

 (3S)-2-Oxo-3-isoquinolin-1-oylamino-5-hydroxyacetyl-N-
- 25 [(2Rs,3s) 2-benzyloxy-5-oxo-tetrahydrofuran-3-yl]2,3,4,5-tetrahydro-7-fluoro-1H-1,5-benzodiazepine-1acetamide(696a-1) was synthesized via methods used to
 prepare 656 to afford 800 as a yellow solid, ¹H NMR
 (500 MHz, CDCl₃) δ 2.55 (ddd, 1H), 2.85 (ddd, 1H),
 30.3,70,3,80 (m. 2H), 3.05 (hm. 2H), 4.05 (dd. 2H), 4.20
- 30 3.70-3.80 (m, 2H), 3.95 (bm, 1H), 4.05 (d, 1H), 4.30 (a, 1H), 4.40-4.60 (m, 4H), 4.70-5.05 (m, 4H), 5.55

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(d, 1H), 7.10 (d, 1H), 7.20-7.35 (m, 3H), 7.40- 7.50 (m, 1H), 7.60- 7.85 (m, 3H), 8.40 (dd, 1H), 9.10 (m, 1H), and 9.30 pp (m, 1H).

(3S)-2-0xo-3-isoquinolin-1-oylamino-5-hydroxyacetyl-N
[(2RS,3S) 2-benzyloxy-5-oxo-tetrahydrofuran-3-yl]2,3,4,5-tetrahydro-7-chloro-1H-1,5-benzodiazepine-1acetamide(696a-2) was synthesized via methods used to
prepare 677, to afford 204 mg of 696a-2 as a white
solid, with the exception that the reduction of the

- nitro- group was done as follows: To a solution of the
 nitro compound (7.2 g, 20 mmol) in MeOH was added NH₄Cl
 (2.1 g, 39 mmol) and Zn (17 g, 260 mmol). The
 resulting mixture was heated to reflux 1 hour after
 which it was cooled and filtered through celite. The
 filtrated was concentrated in vacuo then treated with
 cold 1N HCl to afford 3.6 g of a pale red solid. ¹H
 NMR(CDCl₃) δ 1.85(s, 1H), 2.45(d, 0.5H), 2.50-2.65(m,
 0.5H), 2.80-2.90(m, 0.5H), 2.90-3.00(m, 0.5H), 3.45(s,
 0.5H), 3.55-3.75(m, 1H), 3.85-4.15(m, 2H), 4.25(d, 1H),
- 20 4.40-4.65(m, 2H), 4.70-4.80(m, 0.5H), 4.85-5.15(m, 3H), 5.40(s, 0.5H), 5.60(d, 0.5H), 7.00(d, 0.5H), 7.15-7.90(m, 12.5H), 8.35-8.45(m, 1H), 9.00-9.10(m, 1H), 9.25-9.40(m, 1H)

(3S)-3-[(3S)-2-0xo-3-isoquinolin-1-oylamino-5-

25 hydroxyacetyl-2,3,4,5-tetrahydro-7-fluoro-1H-1,5-benzodiazepine-1-acetylamino]4-oxobutyric acid(696-1) was synthesized via methods used to prepare 2002 from 2001 to afford 140 mg of 696-1 as a white solid, ¹H NMR (500 MHz, CD₃OD) 0 2.50 (m, 1H), 2.70 (m, 1H), 3.85 (d, 30 1H), 3.95 (m, 1H), 4.10 (m, 1H), 4.20 (m, 1H), 3.85 (d, 30 1H), 3.95 (m, 1H), 4.10 (m, 1H), 4.20 (m,

30 1H), 3.95 (m, 1H), 4.10 (d, 1H), 4.35 (m, 1H), 4.50-4.60 (m, 2H), 4.80 (bm, 1H), 5.00 (m, 1H), 7.40-7.48

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(m, 3H), 7.65 (m, 1H), 7.75 (t, 1H), 7.85 (t, 1H), 8.00 (d, 1H), 8.55 (d, 1H), and 9.05 ppm (d, 1H).

(3S)-3-[(3S)-2-Oxo-3-isoquinolin-1-oylamino-5-hydroxyacetyl-2,3,4,5-tetrahydro-7-chloro-1H-1,5
benzodiazepine-1-acetylamino]4-oxobutyric acid(696-2)
was synthesized via methods used to prepare 2002 from 2001 to afford 250 mg of 696-2as a white solid, ¹H
NMR(CD₃OD) & 2.40-2.55 (m, 1H), 2.60-2.75 (m, 1H), 3.80-4.00 (m, 2H), 4.05 (d, 1H), 4.20-4.35 (m, 1H), 4.45
10 4.65 (m, 3H), 4.80-5.10 (m, 2H)

(3S)-2-Oxo-3-isoquinolin-1-oylamino-5-methoxyacetyl-N-[(2RS,3S) 2-benzyloxy-5-oxo-tetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-7-fluoro-1H-1,5-benzodiazepine-1-acetamide(699a-1) was synthesized via methods used to prepare 655 to afford 699a-1 h NMR (500 MHz, CDCl₃) 8 2.55 (ddd, 1H), 2.90 (ddd, 1H), 3.25 (s, 3H), 3.28 (s, 3H), 3.80 (bt, 2H), 3.95 (bm, 2H), 4.25 (dd, 1H), 4.45-4.90 (m, 3H), 5.60 (d, 1H), 7.05-7.40 (m, 8H), 7.50 (bm, 1H), 7.65-7.85 (m, 2H), 8.45 (d, 1H), 9.1 (m,

(3s)-3-[(3s)-2-0xo-3-isoquinolin-1-oylamino-5-methoxyacety1-2,3,4,5-tetrahydro-7-fluoro-1H-1,5-benzodiazepine-1-acetylamino]4-oxobutyric acid(699a-2) was synthesized via methods used to prepare 2002 from 25 2001 to afford 699a-2 ¹H NMR (500 MHz, CD₃OD) & 2.51 (m, 1H), 2.70 (dt, 1H), 3.31 (bs, 3H), 3.90 (bdt, 1H), 3.95 (bm, 1H), 4.05 (d, 1H), 4.35 (m, 1H), 4.50 (d, 1H), 4.60 (dd, 1H), 4.65 (dt, 1H), 4.80 (m, 1H], 5.05 (m, 1H), 7.35- 7.48 (m, 3H], 7.65 (bm, 1H), 7.75 (t,

1H), and 9.35 ppm (m, 1H)

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1H), 7.82 (t, 1H), 8.05 (d, 1H), 8.55 (d, 1H), and 9.05 ppm (d, 1H).

(3S) -3-[(3S) -2-0xo-3-(3,5-dimethyl-4-hydroxybenzoyl) amino-5-methoxyacetyl-2,3,4,5-

- 5 tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4oxobutyric acid, O-2,6-dichlorobenzyl oxime(688c) was
 synthesized via methods used to prepare 308d to afford
 800, ¹H NMR (CD₃OD) δ 2.2 (s, 6H), 2.58-2.83 (m, 2H),
 3.28 (s, 3H), 3.29-3.34 (m, 1H), 3.68-3.80 (m, 2H),
- 10 3.95-4.05 (dd, 1H), 4.38-4.48 (dd, 1H), 4.82-5.00 (m, 2H), 5.26-5.36 (m, 2H), 7.22-7.65 (m, 10H).
 - (3S)-2-Oxo-(2,4-dimethylthiazo-5-yl)amino-5hydroxyacetyl-N-((2RS,3S) 2-benzyloxy-5-oxotetrahydrofuran-3-yl]-2,3,4,5-tetrahydro-1H-1,5-
- 15 benzodiazepine-1-acetamide(800) was synthesized via methods used to prepare 696a-1 to afford 204 mg of 800 as a yellow solid, ¹H NMR(CDCl₃) (mixture of diastereomers) 8 1.70(s, 1H), 2.40-2.80(m, 7H), 2.80- 2.90(m, 0.5H), 2.95-3.05(m, 0.5H), 3.30-3.35(m, 0.5H), 20 3.45-3.55(m, 0.5H), 3.55-3.65(m, 1H), 3.80-4.05(m, 2H),
- 20 3.45-3.55(m, 0.5H), 3.55-3.65(m, 1H), 3.80-4.05(m, 2H) 4.30-4.50(m, 2H), 4.55-4.65(m, 1H), 4.75-4.95(m, 3H), 5.45(s, 0.5H), 5.55(d, 0.5H), 6.70(d, 0.5H), 6.90(d, 0.5H), 7.15-7.80(m, 10H)
 - (3S) -3-[(3S) -2-0xo-3-(2,4-dimethylthiazo-1-oyl)amino-5-
- 25 hydroxyacetyl-2,3,4,5-tetrahydro-1H-1,5-benzodiazepine-1-acetylamino]4-oxobutyric acid(801) was synthesized via methods used to prepare 2002 from 2001 to afford 801.

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Example 34

Compounds 720-73 were prepared by methods similar to the methods used to prepare compounds 619-635 (see, Example 13). Physical data for compounds 5 720-73 is listed in Table 29.

 /41	-

Structure Structure	MF MW HELC RT min min Purity Purity 546.93 10.729 998
	—————————————————————————————————————

MS (M+Na)+	578.2	564.5
HPLC RT min Purity	761 99%	.655 79%
MW	554.56 11.761	540.53 10.655
MF	C27H30N409	С26Н28N409
Structure	H ₂ C C C C C C C C C C C C C C C C C C C	
Compound	722	723

	T	
MS (M+Na)+	563.1	5.77.2
r r t<	on ⊙	90 60
HPLC RT min Purity	538.56 10.584	552.59 11.329 99%
MM	538,56	552,59
MF	C27H30N408	C28H32N408
Structure		
Compound	724	725

		T
MS (M+Na)+	620.8	506.6
7 % t	ა დ გ	928
HPLC RT min Purity	10.667	9.085
MM	596.60 10.667	482.50 9.085
MF	C29H32N4O10	C24H26N4O7
Structure	H ₂ C () 0 0H	
Compound	726	727

	1
634.9	607.3
95	ත. න
11.5	11.611
610.63	582.57 11.611
C30H34N4O10	C28H30N4O1C
H ₂ C Ch ₂ C Ch ₃ C Ch ₃ C Ch ₄ C Ch	
728	729
	H ₂ C C C C C C C C C C C C C C C C C C C

MS (M+Na)+	572.2	5. 1.	
RT n tv	896	928	
HPLC RT min Purity	3.939	4.298	
MW	549.50 3.939 96%	563.53 4.298	
MF	C23H27N5O11	C24H29N5O11	
Structure	How I have been a second and the sec		
Compound	730	731	

Compound	Structure	MF	MM	HPLC RT min Purity	MS (M+Na)+
732		C26H28N4011	572.53	7.640 98%	595.9
733		C25H26N4C10 542.51	542.51	7.375 98%	565.9

PCT/US96/20843

MS (M+Na)+	630.6	632.1	
RT t	96 60 60	92.28	
HPLC RT min Purity	9.656	10.887	
MM	608.62	609.62 10.887	
MF	C32H28N607	C28H27N509S	
Structure			
Compound	734	7.85 2.85	

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Example 35

Compounds 736-767 were prepared by methods similar to the methods used to prepare compounds 619-635 (see, Example 13). Physical data for compounds 5 736-767 is listed in Table 30.

Table 30

Compound	R ⁴	R ³
736		Э
737		HO ====================================
738	H,N-(-)	HO
739	NH H ₃ C CH ₃	HO.
740	O H	HO E
741	an i	но

Compound	R ⁴	R ³
742	NJ i	HOJ
743	i soi	но
744		2
745		0
746	H	10 1 0
747	430-F4	но Ц
748	OH OH	ю
749	HOOLINGOH	ноЩ
750	HOLN	HOJ
751	HOTO	но∬
752	CI NOCH3	HO_I

5

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Compound	R ⁴	R ³
753	H ₃ C	HO
754		ю
755	J. P.	HO.
756	H600 CH 0	ю
757	CH CH	PO={
758	2	₩
759	H _C C H N N N N N N N N N N N N N N N N N N	9 £
760	HC HC HC	0= /
761	CN OH	9 9
762	HO CH	ю
763	N. O. N.	юЩ

5

Compound	R ⁴	R ³
764	H ₃ C NH O	HO.
765		D = 0
766	HN CO	4
767	NY OH O	ю

- 5 The data of the examples above demonstrate that compounds according to this invention display inhibitory activity towards IL-1B Converting Enzyme.

 Insofar as the compounds of this invention are able to inhibit ICE in vitro and furthermore, may be delivered orally to mammals, they are of evident clinical utility for the treatment of IL-1-, apoptosis-, IGIF-, and IFN-v mediated diseases. These tests are predictive of the compounds ability to
- 15 While we have described a number of embodiments of this invention, it is apparent that our basic constructions may be altered to provide other embodiments which utilize the products and processes of this invention. Therefore, it will be appreciated that the scope of this invention is to be defined by the appended claims, rather than by the specific embodiments which have been presented by way of example.

inhibit ICE in vivo.

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CLAIMS

We claim:

1. A compound represented by the formula:

5 $\alpha \qquad \qquad \begin{array}{c} \text{(CJ}_2)_{\pi}\text{-T} \\ \text{(CH}_2)_g\text{-R}_3 \end{array}$

wherein:

10 X₁ is -CH;

15

g is 0 or 1;

each J is independently selected from the group consisting of -H, -OH, and -F, provided that when a first and second J are bound to a C and said first J is -OH, said second J is -H;

m is 0, 1, or 2;

T is -OH, -CO-CO $_2$ H, -CO $_2$ H, or any bioisosteric replacement for -CO $_2$ H;

 R_1 is selected from the group consisting of the following formulae, in which any ring may optionally be singly or multiply substituted at any carbon by Q_1 , at any nitrogen by R_5 , or at any atom by =0, -OH, -CO₂H, or halogen; and any saturated ring may optionally be unsaturated at one or two bonds:

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wherein each ring C is independently chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

$$\begin{array}{c} R_3 \text{ is:} \\ -\text{CN,} \\ -\text{CN-CH-Rp,} \\ 10 & -\text{CH=N-O-Rp,} \\ -(\text{CH}_2)_{1-3} - T_1 - \text{Rp,} \\ -\text{CJ}_2 - \text{Rp,} \\ -\text{CO-R}_{13}, \text{ or} \\ / \text{Ks} \\ 15 & -\text{CO-CO-N} \\ & \text{$\backslash R_{10}$;} \end{array}$$

5

each $\ensuremath{R_4}$ is independently selected from the group consisting of:

 $\mbox{ each T_1 is independently selected from the group} \\ \mbox{25} \qquad \mbox{consisting of:}$

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- 756 -

15

20

30

each R_9 is a C_{1-6} straight or branched alkyl group optionally singly or multiply substituted with -OH, -F, or =0 and optionally substituted with one or two Ar_1 groups;

each ${\rm R}_{10}$ is independently selected from the group consisting of -H or a ${\rm C}_{1-6}$ straight or branched alkyl group;

each $\rm R_{13}$ is independently selected from the group consisting of -Ar_2, -R_4 and -N-OH $$\backslash$$.

each Ar₁ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, a cycloalkyl group which contains between 3 and 15 carbon atoms and between 1 and 3 rings, said cycloalkyl group being optionally benzofused, and a heterocycle group containing between 5 and 15 ring

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atoms and between 1 and 3 rings, said heterocycle group containing at least one heteroatom group selected from -O-, -S-, -SO-, $-\text{SO}_2$ -, =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted with $-\text{NH}_2$, $-\text{CO}_2\text{H}$, -Cl, -F, -Br, -I, $-\text{NO}_2$, -CN,

=0, -OH, -perfluoro
$$C_{1-3}$$
 alkyl, C_{1-2} or C_{1-3} or C_{1-3}

each Ar_2 is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ and $-Q_2$:

$$(jj)$$
 $\longrightarrow_{X-Y}^{N}$; and

20

5

10

15

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each \mathbf{Q}_1 is independently selected from the group consisting of:

$$-Ar_1$$

-R₉,

$$-T_1-R_9$$
, and $-(CH_2)_{1,2,3}-T_1-R_9$;

each Q_2 is independently selected from the group consisting of -OH, -NH₂, -CO₂H, -Cl, -F, -Br, -I,

15

5

provided that when -Ar $_1$ is substituted with a Q_1 group which comprises one or more additional -Ar $_1$ groups, said additional -Ar $_1$ groups are not substituted with Q_1 ;

20 each X is independently selected from the group
consisting of =N-, and =CH-;

each $\rm X_2$ is independently selected from the group consisting of -O-, -CH2-, -NH-, -S-, -SO-, and -SO2-;

 $\mbox{ each Y is independently selected from the group} \\ \mbox{25} \qquad \mbox{consisting of -O-, -S-, and -NH;} \\$

provided that when

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```
X2 is 0,
                    R_5 is benzyloxycarbonyl, and
                    ring C is benzo,
              then R3 cannot be -CO-R13 when:
                    R_{13} is -CH_2-O-Ar_1 and
  5
                    Ar<sub>1</sub> is 1-phenyl-3-trifluoromethyl-
        pyrazole-5-yl wherein the phenyl is optionally
        substituted with a chlorine atom;
              or when
1.0
                    R<sub>13</sub> is -CH<sub>2</sub>-O-CO-Ar<sub>1</sub>, wherein
                    Ar<sub>1</sub> is 2,6-dichlorophenyl.
                    2. The compound according to claim 1,
        wherein:
             X_1 is -CH;
15
              a is 0;
              J is -H:
             m is 0 or 1 and T is -\text{CO-CO}_2\text{H}, or any bioisosteric
        replacement for -CO2H, or
20
             m is 1 and T is -COoH;
             ring C is benze optionally substituted with
       -C_{1-3} alkyl, -O-C_{1-3} alkyl, -Cl, -F or -CF_3;
             Rs is:
                   -CO-Ar<sub>1</sub>
25
                   -SO2-Ar1,
                   -CO-NH2
                   -CO-NH-Ar,
                   -CO-Rg,
```

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 R_7 is -H and R_6 is: -H, -R₉, or -Ar₁;

 R_9 is a C_{1-6} straight or branched alkyl group optionally substituted with =O and optionally substituted with -Arı;

10 R_{10} is H or a $-C_{1-3}$ straight or branched alkyl group;

Ar_1 is phenyl, naphthyl, pyridyl, benzothiazolyl, thienyl, benzothienyl, benzoxazolyl, 2-indanyl, or indolyl optionally substituted with -O-C₁₋₃ alkyl, -NH-C₁₋₃ alkyl, -N-(C₁₋₃ alkyl)₂, -Cl, -F, -CF₃,

 $-C_{1-3}$ alkyl, or

O / \ CH₂ ;

20

25

15

5

 $\rm Q_1$ is $\rm R_9$ or $\rm -(CH_2)_{\,0\,,\,1\,,\,2}-T_1-(CH_2)_{\,0\,,\,1\,,\,2}-Ar_1,$ wherein $\rm T_1$ is -O- or -S-;

each X is independently selected from the group consisting of =N-, and =CH-;

each X_2 is independently selected from the group consisting of -O-, -CH₂-, -NH-, -S-, -SO-, and -SO₂-.

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 $\label{eq:compound} \mbox{3. The compound according to claims 1 or 2,} \\ \mbox{wherein the R_1 group is:}$

(w1) $\begin{array}{c} X_2 \\ \\ R_6 \\ \\ H \\ \end{array}$

5 X_2 is:

-O- ,

-SO₂-, or

-302-, 0.

-NH-;

10

optionally substituted with ${\rm R}_5$ or ${\rm Q}_1$ at ${\rm X}_2$ when ${\rm X}_2$ is -NH-; and

ring C is benzo substituted with $-{\rm C}_{1-3}$ alkyl, $-{\rm O}-{\rm C}_{1-3}$ alkyl, $-{\rm Cl}_1$ -F or $-{\rm CF}_3$.

4. A compound represented by the formula:

(<u>I</u>) R₁-N-R

wherein:

20 R_1 is selected from the group consisting of the following formulae:

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$$(z) \qquad \qquad \begin{matrix} y_2 \\ y_2 \\ y_3 \\ y_4 \\ y_5 \\ y_6 \\ y_7 \\ y_8 \\ y_9 \\$$

ring C is chosen from the group consisting of
benzo, pyrido, thieno, pyrrolo, furano, thiazolo,
5 isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo,
cyclopentyl, and cyclohexyl;

R2 is:

10 m is 1 or 2;

 R_5 is selected from the group consisting of: $\label{eq:constant} -\text{C(O)-R}_{10}, \\ -\text{C(O)-R}_{9},$

20
$$-S(O)_2-R_9$$
, $-C(O)-CH_2-O-R_9$,

$$-C(0)C(0)-R_{10}$$
,
 $-R_{9}$,
 $-H$, and
 $-C(0)C(0)-OR_{10}$;

5
$$X_5$$
 is -CH- or -N-; X_5 is X_5

$$X_7$$
 is $-N(R_8)$ - or $-O-$;

10

30

 $\rm R_{6}$ is selected from the group consisting of -H and -CH $_{\rm 3};$

 $\ensuremath{R_8}$ is selected from the group consisting of:

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

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each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -Cl₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -Cl₁₋₆ alkyl group is optionally unsaturated;

 $\rm R_{13}$ is selected from the group consisting of H, Ar₃, and a C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

each R_{51} is independently selected from the group consisting of R_9 , $-C(0)-R_9$, $-C(0)-N(H)-R_9$, or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

each R_{21} is independently selected from the group consisting of -H or a -C₁₋₆ straight or branched alkyl group;

15

20

25

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₂;

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each Q_1 is independently selected from the group consisting of -NH $_2$, -CO $_2$ H, -Cl, -F, -Br, -I, -NO $_2$, -CN, =O, -OH, -perfluoro C $_{1-3}$ alkyl, R $_5$, -OR $_5$, -NHR $_5$, -OR $_9$, -NHR $_9$, -R $_9$, -C(O)-R $_{10}$, and

5

10

provided that when $-Ar_3$ is substituted with a O_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

15 5. The compound according to claim 4, wherein R_5 is selected from the group consisting of:

$$-C(0)-R_{10}$$
,
 $-C(0)O-R_{9}$, and

20 6. The compound according to claim 4, wherein R_5 is selected from the group consisting of:

$$-R_9$$
, and

7. The compound according to claims 5 or $\boldsymbol{\varepsilon},$ wherein:

m is 1;

25

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with $-A_{T_3}$, -OH, $-OR_9$, $-CO_2H$, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein A_{T_3} is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

$$R_{21}$$
 is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group optionally substituted with $-Ar_3$, wherein Ar_3 is phenyl, optionally substituted by $-O_1$;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₅, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₄, and

wherein each R_9 and R_{10} are independently a $-C_{1-\ell}$ straight or branched alkyl group optionally substituted with $-Ar_1$ wherein Ar_2 is phenyl;

3.0

20

25

5

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provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional -Ara groups, said additional -Ar3 groups are not substituted with another -Ar3.

5 8. A compound represented by the formula:

wherein:

m is 1 or 2;

10 R_1 is selected from the group consisting of the following formulae:

(e10)

, wherein X5 is N;

(e11) 15

(e12)

(w2)

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5

10

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

 R_3 is selected from the group consisting of: -CN, -C(O)-H, -C(O)-CH₂-T₁-R₁₁, -C(O)-CH₂-F, -C=N-O-R₉, and -CO-Ar₂;

 $$R_{5}$$ is selected from the group consisting of: $\label{eq:constraint} -\text{C(O)-R}_{10},$ 20 $-\text{C(O)O-R}_{9},$

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-C(0)-C(0)-OR₁₀;

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each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated; .

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

10 each ${\rm R}_{11}$ is independently selected from the group consisting of:

-Ar₄,
- (CH₂)₁₋₃-Ar₄,
-H, and
-C(0)-Ar₄;

15

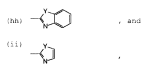
 $\rm R_{13}$ is selected from the group consisting of H, Ar₃, and a C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

20 OR₁₃ is optionally -N(H)-OH;

each R_{21} is independently selected from the group consisting of -H or a $-C_{1-6}$ straight or branched alkyl group;

Ar $_2$ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by Q_1 :

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1.0

15

20

25

wherein each Y is independently selected from the group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

each Ar_4 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q1;

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each Q_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =O, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, -OR₉, -NHR₉, -R₉, -C(O)-R₁₀, and

5

10

provided that when -Ar $_3$ is substituted with a Q $_1$ group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

- 9. The compound according to claim 8, wherein R_1 is (e11).
 - 10. The compound according to claim 8, wherein $\ensuremath{R_1}$ is (e12).
- $\label{eq:compound} 11. \quad \text{The compound according to claim θ,}$ 20 $\quad \text{wherein R_3 is (y1).}$
 - 12. The compound according to claim 8, wherein \boldsymbol{R}_1 is $(\boldsymbol{y}2)$.
 - $\label{eq:compound} 13. \quad \text{The compound according to claim 8,} \\ \text{wherein and } R_1 \text{ is (z).}$
- - 15. The compound according to claim 14, wherein:

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m is 1;

ring C is benzo, pyrido, or thieno;

 R_3 is selected from the group consisting of -C(0)-H, -C(0)-Ar_2, and -C(0)CH_2-T_1-R_1;

 $R_5 \ \mbox{is selected from the group consisting of:} \\ -C(0)-R_{10}, \ \mbox{wherein } R_{10} \ \mbox{is } -Ar_3; \\ -C(0)0-R_{9}, \ \mbox{wherein } R_9 \ \mbox{is } -CH_2-Ar_3; \\ -C(0)C(0)-R_{10}, \ \mbox{wherein } R_{10} \ \mbox{is } -Ar_3; \\ \end{array}$

-R₉, wherein R₉ is a C_{1-2} alkyl group substituted with -Ar₃; and -C(O)C(O)-OR₁₀, wherein R₁₀ is -CH₂Ar₃;

T₁ is 0 or S;

Re is H;

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15 $R_{B} \text{ is selected from the group consisting } -C(0)-R_{10}, \\ -C(0)-CH_{2}-OR_{10}, \text{ and } -C(0)CH_{2}-N(R_{10})(R_{10}), \text{ wherein } R_{10} \text{ is} \\ H, CH_{3}, \text{ or } -CH_{2}CH_{3};$

 $\rm R_{11}$ is selected from the group consisting of -Ar4, -(CH2) $_{1-3}\text{-Ar}_4$, and -C(O)-Ar4;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, -CO₂H, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O_1 ;

25 Ar₂ is (hh);

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Y is O:

5

2.0

3.0

each Ar_3 cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl, thienothienyl, thiadiazolyl, benzoftriazolyl, benzo[b]thiophenyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -0_1 ;

each Ar₄ cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, naphthyl, pyridinyl, oxazolyl, pyrimidinyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -01;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -OR₉, -NHR₉, and

O /\ CH₂,

wherein each R_9 and R_{10} are independently a $-C_{1-\epsilon}$ 25 straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_2$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

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. 16. The compound according to claim 8, wherein R_1 is (e10) and X_5 is N.

- $$17.$\,$ The compound according to claim 16, wherein R_{3} is CO-Ar $\!_{2}.$
- - $\begin{tabular}{ll} 19. & The compound according to claim 16, \\ wherein: \\ \end{tabular}$

$$R_3$$
 is $-C(0)-CH_2-T_1-R_{11}$;
 T_1 is O ; and
 R_{11} is $-C(0)-Ar_4$.

- 20. The compound according to claim 16, wherein $R_{\rm 3}$ is -C(O)-H.
- $\label{eq:21.2} 21. \quad The compound according to claim 16, \\ 15 \qquad \mbox{wherein R_3 is $-CO-CH_2-T_1-R_{11}$ and R_{11} is $-Ar_4$.}$
 - 22. The compound according to any one of claims 19-21, wherein $R_{\hat{S}}$ is selected from the group consisting of:

$$-C(0)-R_{10}$$
,
20 $-C(0)O-R_{9}$, and $-C(0)-NH-R_{10}$.

\$23.\$ The compound according to claim 22, wherein:

m is 1;

25

T, is O or S,

- 777 ~

provided that when ${\rm R}_3$ is -C(0)-CH₂-T₁-R₁₁, T₁ is C:

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, -CO₂H, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with O_{13}

 R_{21} is -H or -CH₃;

 Ar_2 is (hh);

10 Y is O;

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁:

each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by -0_1 ;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₋₀, -OR₉, -NHR₉, and

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

 $24.\$ The compound according to any one of claims 19-21, wherein R_5 is selected from the group consisting of:

-S(O)2-Rg,

-S(O)2-NH-R10,

-C(0)-C(0)-R10,

-Rg, and

-C(0)-C(0)-OR₁₀.

\$25.\$ The compound according to claim 24, wherein:

m is 1;

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 T_1 is 0 or S,

provided that when R_3 is -C(0)-CH $_2$ -T $_1$ -R $_{11},\ T_1$ is 0;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, $-CO_7H$,

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wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

$$R_{21}$$
 is -H or -CH₃;

5 Ar₂ is (hh);

10

1.5

20

25

Y is 0;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thiencthienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -0:

each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₉, and

- 780 -

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_2$ wherein Ar_2 is phenyl;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

26. A compound represented by the formula:

wherein:

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R₁ is:

R₅ is selected from the group consisting of:

-C(O)O-Ra,

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- 781 -

-S(0)₂-R₉, -C(0)-CH₂-O-R₉, -C(0)C(0)-R₁₀, -R₉, -H, and -C(0)C(0)-OR₁₀.

Xs is CH;

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Y2 is H2 or O;

each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with -Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalxyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 $\rm R_{13}$ is selected from the group consisting of H, Ar₃, and a C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

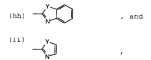
OR₁₃ is optionally -N(H)-OH;

each R_{21} is independently selected from the group consisting of -H or a $-C_{1-6}$ straight or branched alkyl group;

Ar₂ is independently selected from the following

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group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by O_1 :



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wherein each Y is independently selected from the group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q-;

each Ar $_4$ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO2, =N-, -NH-, -N(Rs)-, and -N(Rg)- said heterocycle group optionally

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containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -0_1 .

each O_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =O, -OH, -perfluoro C₁₋₃ alkyl, R₅, -OR₅, -NHR₅, -OR₉, -NHR₉, -R₉, -C(O)-R₁₀, and

O / \ CH₂;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

27. A compound represented by the formula:

wherein:

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1.0

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m is 1 or 2:

R; is:

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$$(e10) \qquad \begin{array}{c} Y_2 \\ R_{21} \\ R_{5} \\ N \end{array} \qquad ; \qquad \qquad \\ \\ R_{5} \\ N \end{array}$$

 R_3 is $-C(0)-CH_2-T_1-R_{11}$ and R_{11} is $-(CH_2)_{1-3}-Ar_4$;

 $\ensuremath{\text{R}}_5$ is selected from the group consisting of:

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-S(O)₂-R₉,

-C(O)-CH2-O-R9,

-C(0)C(0)-R₁₀

-C(0)C(0)-R₁₀

-R₉, -H. and

-C(0)C(0)-OR₁₀

Xs is CH;

Y2 is H2 or O;

each T_1 is independently selected from the group consisting of -0-, -S-, -S(0)-, and -S(0)₂-;

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group

- 785 -

consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 $\rm R_{13}$ is selected from the group consisting of H, Ar₃, and a C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

OR₁₃ is optionally -N(H)-OH;

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each R_{21} is independently selected from the group consisting of -H or a -C₁₋₆ straight or branched alkyl group;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

each Ar₄ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said

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heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =0, -OH, -perfluoro C_{1-3} alkyl, -R₅, -OR₅, -NHR₅, -OR₉, -NHR₀, -R₉, -C(O)-R₁₀, and

provided that when -Ar $_3$ is substituted with a \mathcal{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

\$28\$. The compound according to claims 26 or 27, wherein $R_{\mbox{\scriptsize 5}}$ is selected from the group consisting of:

25
$$-C(0)-R_{10}$$
, $-C(0)0-R_{9}$, and $-C(0)-NH-R_{10}$.

\$29.\$ The compound according to claim 28, wherein:

30 m is 1;

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 T_1 is 0 or S;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, $-CO_2H$, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with C_1 ;

 R_{21} is -H or -CH₃;

Ar2 is (hh);

Y is O:

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -O₁:

each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₉, and

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wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_2$ wherein Ar_2 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

30. The compound according to claims 26 or 15 27, wherein $R_{\bar{5}}$ is selected from the group consisting of:

 $-S(0)_2-R_9$, $-S(0)_2-NH-R_{10}$,

-C(O)-C(O)-R₁₀,

-R₉, and

-C(O)-C(O)-OR₁₀.

 $$31.$\,$ The compound according to claim 30, wherein:

m is 1;

25

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 T_1 is 0 or S;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, $-CO_0H$, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar_3 is morpholinyl or phenyl,

- 789 -

wherein the phenyl is optionally substituted with Q_1 ;

 R_{21} is -H or -CH₃;

Aro is (hh);

Y is O;

5

1.0

15

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo(b)thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -01;

each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₉, and

О / \ СН \ /

wherein each ${\rm R}_9$ and ${\rm R}_{10}$ are independently a $-{\rm C}_{1-6}$ straight or branched alkyl group optionally substituted

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with -Ar3 wherein Ar3 is phenyl;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

32. A compound represented by the formula:

wherein:

5

R₁ is:

(e10)

 R_3 is -C(0)-CH₂-T₁-R₁₁; T_1 is 0; and R_{11} is

15 -C(O)-Ar4;

20

 $\ensuremath{\mathtt{R}}_5$ is selected from the group consisting of:

-S(0)2-Rg,

-S(0)2-NH-R10,

-C(0)-C(0)-R₁₀,

-Rg, and

-C(0)-C(0)-OR10;

Xs is CH;

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Y2 is H2 or O;

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each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with -Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 R_{13} is selected from the group consisting of H, Ar₃, and a C_{1-6} straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

15 OR₁₃ is optionally -N(H)-OH;

each $\rm R_{21}$ is independently selected from the group consisting of -H or a -C_{1-6} straight or branched alkyl group;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings,

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and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each Ar_4 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =O, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, -OR₉, -NHR₉, -R₉, -C(O)-R₁₀, and

O / \ CH₂;

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provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

33. A compound represented by the formula:

wherein:

m is 1 or 2;

R₁ is:

5 (e10)

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 R_3 is -C(0)-H;

 R_5 is selected from the group consisting of:

;

-S(O)2-Rg,

-S(O)2-NH-R10,

-C(0)-C(0)-R₁₀,

-Rg, and

-C(0)-C(0)-OR10;

15 X₅ is CH;

 Y_2 is H_2 or O;

each R_{9} is independently selected from the group consisting of $-Ar_{3}$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_{3}$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

each $R_{\rm 1C}$ is independently selected from the group consisting of -H, -Ar_3, a -C_{3-6} cycloalkyl group, and a

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 $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

 R_{13} is selected from the group consisting of H, Ar₃, and a C_{1-6} straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

OR13 is optionally -N(H)-OH;

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each R_{21} is independently selected from the group consisting of -H or a $-C_{1-6}$ straight or branched alkyl group;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one hetercatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q1;

each Q₁ is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, -C, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, -OR₉, -NHR₉, -R₉, -C(0)-R₁₀, and

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provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

34. The compound according to claims 32 or 33, wherein:

m 1s 1:

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15 R₁₃ is H or a C_{1-4} straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, -CO₂H, wherein the R₉ is a C_{1-4} branched or straight chain alkyl group; wherein Ar₃ is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

20 R₂₁ is -H or -CH₃;

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁;

30 each Ar₄ cyclic group is independently selected from the set consisting of phenyl, tetrazolyl,

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pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by $-O_1$:

each Q_1 is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₉, and

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

35. A compound represented by the formula:

wherein:

m is 1;

25 R₁ is:

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- 797 -

$$\begin{array}{c} \text{(el0)} \\ & \text{R}_{2}\text{-} \\ & \text{R}_{5}\text{-} \\ & \text{H} \end{array} ;$$

 R_3 is -CO-CH₂-T₁-R₁₁ and R_{11} is -Ar₄;

 R_5 is selected from the group consisting of: $-C(0)-R_{10}$, $-C(0)O-R_9$, and $-C(0)-NH-R_{10}$;

X5 is CH;

Y2 is 0;

10 T₁ is O or S;

5

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each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with -Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with -Ar₃, -OH, -OR₉, -CO₂H, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar₃ is morpholinyl or phenyl,

- 798 -

wherein the phenyl is optionally substituted with Q_1 ;

R21 is -H or -CH3;

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo(b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -O:;

each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by -0_1 ;

each Q₂ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -OR₉, -NHR₉, and

CH₂,

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$

- 799 -

groups, said additional $-\mathrm{Ar}_3$ groups are not substituted with another $-\mathrm{Ar}_3$.

36. A compound represented by the formula:

5 wherein:

m is 1;

R₁ is:

(e10)

15

 R_3 is $-CO-CH_2-T_1-R_{11}$ and R_{11} is $-Ar_4$;

R₅ is selected from the group consisting of:

-S(O)2-Rg,

 $-S(0)_2-NH-R_{10}$

-C(0)-C(0)-R₁₀,

 $-R_g$, and

-C(0)-C(0)-OR10;

X5 is CH;

Y2 is 0;

20 T₁ is 0 or S;

- 800 -

each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with -Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with $-Ar_3$, -OH, $-OR_9$, $-CO_2H$, wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted with Q_1 ;

15 R₂₁ is -H or -CH₃;

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each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by -0.;

each Ar_4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

- 801 -

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(O)-R₁₀ or -S(O)₂-R₉, -OR₅ wherein R₅ is -C(O)-R₁₀, -NHR₀, and

5 O CH₂

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

\$ 37. The compound according to claim 7 selected from the group consisting of:

20 213e

302 CH

- 802 -

5 38. The compound according to claims θ or $_{\rm 68},$ selected from the group consisting of:

- 803 -

- 804 -

- 805 -

5 39. The compound according to claim 15 selected from the group consisting of:

- 806 -

- 807 -

- 808 -

- 809 -

- 811 -

- 813 -

40. The compound according to claims 8 or 68, selected from the group consisting of:

217c

- 814 -

281	H OO H BF/ O	;
282		;
283		;
284	H,CO L C C C C C C C C C C C C C C C C C C	;
285	H ₂ CO H O CH ₃	;
286	H _{SCO} H O O	;

- 816 -

- 817 -

- 818 -

- 820 -

- 821 -

- 822 -

- 823 -

- 824 -

- 825 -

- 826 -

- 828 -

5 477 ;

- 830 -

- 831 -

- 832 -

- 833 -

- 834 -

- 836 -

1007

1008 , H OH ,

1009 ;

1010 ;

1011

, NO OH

- 838 -

1025

1026

5 1030

- 840 -

1035

- 841 -

1037

1038

1039

1040

1041

5

NH OH H

H₅C N H O N H

- 842 -

1042 ;

1043

1044 ;

1045 ,

5 1046 ;

- 843 -

- 844 -

- 846 -

- 848 -

- 849 -

- 853 -

5 41. The compound according to claim 33 selected from the group consisting of:

- 854 -

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42. A pharmaceutical composition comprising

- 855 -

an ICE inhibitor according to any one of claims 1-41 and 57-135 in an amount effective for treating or preventing an IL-1-mediated disease and a pharmaceutically acceptable carrier.

- 5 43. A pharmaceutical composition comprising an ICE inhibitor according to any one of claims 1-41 and 57-135 in an amount effective for treating or preventing an apoptosis-mediated disease and a pharmaceutically acceptable carrier.
- 44. The pharmaceutical composition according to claim 42, wherein the IL-1-mediated disease is an inflammatory disease selected from the group consisting of osteoarthritis, acute pancreatitis, chronic pancreatitis, asthma, and adult respiratory distress syndrome.
 - 45. The pharmaceutical composition according to claim 44, wherein the inflammatory disease is osteoarthritis or acute pancreatitis.

46. The pharmaceutical composition according

20 to claim 42, wherein the IL-1-mediated disease is an autoimmune disease selected from the group consisting of glomeralonephritis, rheumatoid arthritis, systemic lupus erythematosus, scleroderma, chronic thyroiditis, Grave's disease, autoimmune gastritis, insulindependent diabetes mellitus (Type I), autoimmune hemolytic anemia, autoimmune neutropenia, thrombocytopenia, chronic active hepatitis, myasthenia gravis, inflammatory bowel disease, Crohn's disease,

psoriasis, and graft vs host disease.

- 856 -

- 47. The pharmaceutical composition according to claim 46, wherein the autoimmune disease is rheumatoid arthritis, inflammatory bowel disease, or Crohn's disease, or psoriasis.
- 5 48. The pharmaceutical composition according to claim 42, wherein the IL-1-mediated disease is a destructive bone disorder selected from the group consisting of osteoporosis or multiple myeloma-related bone disorder.
- 49. The pharmaceutical composition according to claim 42, wherein the IL-1-mediated disease is a proliferative disorder selected from the group consisting of acute myelogenous leukemia, chronic myelogenous leukemia, metastatic melanoma, Kaposi's sarcoma, and multiple myeloma.
 - 50. The pharmaceutical composition according to claim 42, wherein the IL-1-mediated disease is an infectious disease, selected from the group consisting of sepsis, septic shock, and Shigellosis.
- 51. The pharmaceutical composition according to claim 42, wherein the IL-1-mediated disease is a degenerative or necrotic disease, selected from the group consisting of Alzheimer's disease, Parkinson's disease, cerebral ischemia, and myocardial ischemia.
- 25 52. The pharmaceutical composition according to claim 51, wherein the degenerative disease is Alzheimer's disease.
 - 53. The pharmaceutical composition according

to claim 43, wherein the apoptosis-mediated disease is a degenerative disease, selected from the group consisting of Alzheimer's disease, Parkinson's disease, cerebral ischemia, myocardial ischemia, spinal muscular atrophy, multiple sclerosis, AIDS-related encephalitis, HIV-related encephalitis, aging, alopecia, and neurological damage due to stroke.

54. A pharmaceutical composition for inhibiting an ICE-mediated function comprising an ICE inhibitor according to any one of claims 1-41 and 57-135 and a pharmaceutically acceptable carrier.

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55. A method for treating or preventing a disease selected from the group consisting of an IL-1 mediated disease, an apoptosis mediated disease, an inflammatory disease, an autoimmune disease, a 15 destructive bone disorder, a proliferative disorder, an infectious disease, a degenerative disease, a necrotic disease, osteoarthritis, pancreatitis, asthma, adult respiratory distress syndrome, glomeralonephritis, 20 rheumatoid arthritis, systemic lupus erythematosus, scleroderma, chronic thyroiditis, Grave's disease, autoimmune gastritis, insulin-dependent diabetes mellitus (Type I), autoimmune hemolytic anemia, autoimmune neutropenia, thrombocytopenia, chronic active hepatitis, myasthenia gravis, inflammatory bowel 25 disease, Crohn's disease, psoriasis, graft vs host disease, osteoporosis, multiple myeloma-related bone disorder, acute myelogenous leukemia, chronic myelogenous leukemia, metastatic melanoma, Kaposi's 30 sarcoma, multiple myeloma, sepsis, septic shock, Shigellosis, Alzheimer's disease, Parkinson's disease, cerebral ischemia, myocardial ischemia, spinal muscular

- 858 -

atrophy, multiple sclerosis, AIDS-related encephalitis, HIV-related encephalitis, aging, alopecia, and neurological damage due to stroke in a patient comprising the step of administering to said patient a pharmaceutical composition according to any one of claims 42 to 54.

- 56. The method according to claim 55, wherein the disease is selected from the group consisting of osteoarthritis, acute pancreatitis, rheumatoid arthritis, inflammatory bowel disease, Crohn's disease, psoriasis, and Alzeheimer's disease.
 - 57. A compound represented by the formula:

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 $\ensuremath{R_{\mathrm{1}}}$ is selected from the group consisting of the following formulae:

wherein:

- 859 -

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl;

Ro is:

m is 1 or 2;

5 each $R_{\rm 5}$ is independently selected from the group consisting of:

$$\text{-C (O) -N }(\text{R}_{10}) \text{ }(\text{R}_{10})$$

10

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$$-C(0)C(0)-OR_{10}$$
, and $-C(0)C(0)-N(R_9)(R_{10})$;

$$X_7$$
 is $-N(R_8)$ - or $-O$ -;

 $$R_{6}$$ is selected from the group consisting of -H and $$^{-\text{CH}_{3}}$;}$

- 861 -

 $\begin{array}{c} R_8 \text{ is selected from the group consisting of:} \\ -C(0)-R_{10}, \\ -C(0)-R_{9}, \\ -C(0)-N(H)-R_{10}, \\ \\ 5 \end{array} \\ \begin{array}{c} -S(0)_2-R_9, \\ -S(0)_2-NH+R_{10}, \\ -C(0)-CH_2-OR_{10}, \\ -C(0)-CH_2N(R_{10}), \\ -C(0)-CH_2N(R_{10}), \\ -C(0)-CH_2C(0)-O-R_9, \\ -C(0)-CH_3C(0)-R_9, \\ -H, \text{ and } \\ -C(0)-C(0)-C(0)-R_{10}; \end{array}$

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

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3.0

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 $\rm R_{13}$ is selected from the group consisting of H, Ar₃, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

each R_{51} is independently selected from the group consisting of R_9 , $-\text{C}(0)-R_9$, $-\text{C}(0)-N(\text{H})-R_9$, or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

- 862 -

each R_{21} is independently selected from the group consisting of -H or a -C₁₋₆ straight or branched alkyl group;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Q₁ is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, -O, -OH, -perfluoro C₁₋₃ alkyl, R₅, -OR₅, -NHR₅, -OR₉, -N(R₉) (R₁₀), -R₉, -C(O)-R₁₀, and O

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provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

\$58.\$ The compound according to claim 57, wherein R_1 is (W2).

59. The compound according to claim 57,

- 863 -

wherein R_1 is (e10) and X_5 is CH.

- \$60.\$ The compound according to claim 57, wherein R_{1} is (e10) and X_{5} is N.
- 61. The compound according to claim 57,5 selected from the group consisting of:

- 864 -

62. A compound represented by the formula:

5 wherein:

m is 1 or 2;

 $\ensuremath{\mathtt{R}}_1$ is selected from the group consisting of the following formulae:

- 865 -

$$(e11) \qquad \qquad \qquad ;$$

$$R_{S} = N \qquad \qquad ;$$

$$(e12) \qquad \qquad \qquad \qquad ;$$

$$R_{S} = N \qquad \qquad ;$$

$$R_{S}$$

ring C is chosen from the group consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo,

15

- 866 -

```
cyclopentyl, and cyclohexyl;
```

```
R_3 is selected from the group consisting of: 
 -CN, 
 -C(0)-H, 
 -C(0)-CH_2-T_1-R_{11}, 
 -C(0)-CH_2-F, 
 -C=N-O-R_9, and 
 -CO-Ar_2;
```

each R_5 is independently selected from the group consisting of:

 $\begin{array}{c} -C(0) - R_{10}, \\ -C(0) - R_{9}, \\ -C(0) - N(R_{10}) (R_{10}) \\ -S(0) _{2} - R_{9}, \\ -S(0) _{2} - NH - R_{10}, \\ -C(0) - CH_{2} - O - R_{9}, \\ -C(0) C(0) - R_{10}, \end{array}$

5

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 $-R_9$, -H, 20 $-C(0)C(0)-0R_{10}$, and $-C(0)C(0)-N(R_9)(R_{10})$;

 Y_2 is H_2 or O;

 X_7 is $-N(R_8)$ - or -O-;

25 each T_1 is independently selected from the group consisting of -O-, -S-, -S(O)-, and -S(O)₂-;

 ${\rm R}_{\rm 6}$ is selected from the group consisting of -H and -CH3;

30 R₈ is selected from the group consisting of:

- 867 -

$$\begin{array}{c} -C(O) - R_{10}, \\ -C(O) - R_{9}, \\ -C(O) - NH - R_{10}, \\ -S(O) _{2} - R_{9}, \\ \end{array}$$

$$\begin{array}{c} -S(O) _{2} - NH - R_{10}, \\ -C(O) - CH_{2} - OR_{10}, \\ -C(O) - CH_{2} - N(R_{10}) (R_{10}), \\ -C(O) - CH_{2} - C(O) - C(O) - CH_{2} - C(O) - C$$

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each R_{11} is independently selected from the group consisting of:

$$-Ar_4$$
,
 $-(CH_2)_{1-3}-Ar_4$,
 $-H$, and
 $-C(0)-Ar_4$;

15

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2.5

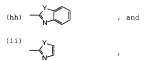
30

 $\rm R_{15}$ is selected from the group consisting of -OH, -OAr_3, -N(H)-OH, and -OCl_-6, wherein $\rm C_{1-6}$ is a straight or branched alkyl group optionally substituted with

- 868 -

each R_{21} is independently selected from the group consisting of -H or a -C $_{1-6}$ straight or branched alkyl group;

Ar₂ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by Q_1 :



1.0

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wherein each Y is independently selected from the group consisting of O and S;

each Ar₃ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Ar4 is a cyclic group independently selected

- 869 -

from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said

5 heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁:

- provided that when $-Ar_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.
- $\mbox{63. The compound according to claim 62,} \\ \mbox{25} \qquad \mbox{wherein } R_1 \mbox{ is } (w2) \, . \\ \mbox{}$

15

- 64. The compound according to claim 62, wherein \mathbf{R}_1 is (el0-A).
 - 65. A compound represented by the formula:

$$(V) \qquad \qquad (I) \qquad R_1 - N \qquad R_3$$

wherein:

(e10-B)

 ${\sf R}_3$ is selected from the group consisting of:

$$-C(0)-CH_2-T_1-R_{11}$$
,

each $\ensuremath{R_5}$ is independently selected from the group

consisting of:

- 871 -

 $-C(0)C(0)-OR_{10}$, and $-C(0)C(0)-N(R_9)(R_{10})$;

 Y_2 is H_2 or O;

5 each T_1 is independently selected from the group consisting of -O-, -S-, -S(0)-, and -S(0)₂-;

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each $\ensuremath{\mathtt{R}}_{11}$ is independently selected from the group consisting of:

-Ar₄, -(CH₂)₁₋₃-Ar₄, -H, and -C(O)-Ar₄;

15

 R_{15} is selected from the group consisting of -OH, -OAr₃, -N(H)-OH, and -OC₁₋₆, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

 R_{21} is $-CH_3$;

 Ar_2 is independently selected from the following

- 872 -

group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by Q_1 :

(hh)
$$\stackrel{\mathsf{Y}}{\longleftarrow}$$
 , and

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wherein each Y is independently selected from the group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁;

each Ar₄ is a cyclic group independently selected from the set consisting of an aryl group which contairs 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, -NH-, -N(Rs)-, and -N(Rs)- said heterocycle group optionally

- 873 -

containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

66. A compound represented by the formula:

wherein:

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20 m is 1 or 2;

25 R_3 is selected from the group consisting cf: -CN,

- 874 -

each R_5 is $-C(0)C(0)-OR_{10}$;

Y2 is H2 or O;

5

15

25

each T_1 is independently selected from the group consisting of -O-, -S-, -S(0)-, and -S(0)₂-;

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

- each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;
- 20 each R_{11} is independently selected from the group consisting of:

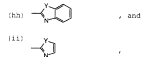
 R_{15} is selected from the group consisting of -OH, -OAr $_3$, -N(H)-OH, and -OC $_{1-6}$, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with

- 875 -

 $-Ar_3$, $-CONH_2$, $-OR_5$, -OH, $-OR_9$, or $-CO_2H$;

each $\rm R_{21}$ is independently selected from the group consisting of -H or a -C1-6 straight or branched alkyl group;

Ar $_2$ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by Q_1 :



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wherein each Y is independently selected from the group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, -N(R_5)-, and -N(R_9)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₁;

each Ar4 is a cyclic group independently selected

- 876 -

from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O.;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

\$67.\$ The compound according to claim 66, wherein R_{21} is $-CH_{3}\,.$

68. A compound represented by the formula:

wherein:

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- 877 -

m is 1 or 2;

R₁ is:

5 (e10-B)

 $\ensuremath{\text{R}}_3$ is selected from the group consisting of:

-CN,

-C(O)-H,

 $-C(0) - CH_2 - T_1 - R_{11}$,

-C(O)-CH2-F,

-C=N-O-Rq, and

-CO-Ar2;

each R5 is independently selected from the group

15 consisting of:

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25

-C(O)-R10,

-C(O)O-R₉,

-C(0)-N(R₁₀)(R₁₀)

-S(O)2-Rg,

-S(O)2-NH-R10,

-C(0)-CH2-O-R9,

-C(0)C(0)-R₁₀,

-Rg.

-H,

 $-C(0)C(0)-OR_{10}$, and

-C(0)C(0)-N(R9)(R10);

Y2 is H2 or O;

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each T_1 is independently selected from the group consisting of -O-, -S-, -S(0)-, and -S(0)₂-;

each R_9 is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched alkyl group optionally substituted with -Ar $_3$, wherein the -C $_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each \mathbf{R}_{11} is independently selected from the group consisting of:

-Ar₄,
- (CH₂)₁₋₃-Ar₄,
-H, and
-C(O)-Ar₄;

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2.0

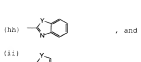
2.5

 R_{15} is selected from the group consisting of -OH, -OAr₃, -N(H)-OH, and -OC₁₋₆, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

each R_{21} is independently selected from the group consisting of -H or a $-C_{1-6}$ straight or branched alkyl group;

Ar $_2$ is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by -Q $_1$ or phenyl, optionally substituted by Q $_1$:

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wherein each Y is independently selected from the group consisting of O and S;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O-;

each Ar₄ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q₁:

- 880 -

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$;

provided that when:

m is 1; R_{15} is -OH; R_{21} is -H; and

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 Y_2 is O and R_3 is -C(O)-H, then R_5 cannot be: -C(O)- R_{10} , wherein R_{10} is -Ar₃ and the Ar₃ cyclic group is phenyl, unsubstituted by - Q_1 , 4- (carboxymethoxy)phenyl, 2-fluorophenyl, 2-pyridyl, N-(4-methylpiperazino)methylphenyl, or

-C(0)-OR9, wherein R9 is -CH2-Ar3, and the Ar3 cyclic group is phenyl, unsubstituted by -Q1; and when

Y₂ is O, R₃ is $-C(0)-CH_2-T_1-R_{11}$, T₁ is O, and R₁₁ is Ar₄, wherein the Ar₄ cyclic group is 5-(1-(4-chlorophenyl)-3-trifluoromethyl)pyrazolyl), then R₅ cannot be:

-H:

-C(0)- R_{10} , wherein R_{10} is -Ar₃ and the Ar₃ cyclic group is 4-(dimethylaminomethyl)phenyl, phenyl, 4-(carboxymethylthio)phenyl,4-(carboxymethylthio)phenyl,

- 881 -

4-(carboxyethyl)phenyl, 4-(carboxypropyl)phenyl, 2-fluorophenyl, 2-pyridyl, N-(4-methylpiperazino)methylphenyl, or

-C(0)-OR9, wherein R9 is isobutyl or -CH2-Ar3 and the Ar3 cyclic group is phenyl;

and when R_{11} is Ar_4 , wherein the Ar_4 cyclic group is 5-(1-phenyl-3-trifluoromethyl)pyrazolyl or 5-(1-(4-chloro-2-pyridinyl)-3-trifluoromethyl)pyrazolyl, then R_5 cannot be:

 $-C(0)-OR_9$, wherein R_9 is $-CH_2-Ar_3$, and the Ar_3 cyclic group is phenyl;

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3.0

and when R_{11} is Ar_4 , wherein the Ar_4 cyclic group is 5-(1-(2-pyridyl)-3-trifluoromethyl)pyrazolyl), then R_5 cannot be:

 $-C(0)-R_{10}$, wherein R_{10} is $-Ar_3$ and the Ar_3 cyclic group is 4-(dimethylaminomethyl) phenyl, or

-C(O)-OR $_9$, wherein R $_9$ is -CH $_2$ -Ar $_3$, and the Ar $_3$ cyclic group is phenyl, unsubstituted by -Q $_1$; and when

 $\rm Y_2$ is O, $\rm R_3$ is -C(O)-CH₂-T₁-R₁₁, T₁ is O, and R₁₁ is -C(O)-Ar₄, wherein the Ar₄ cyclic group is 2,5-dichlorophenyl, then R₅ cannot be:

-C(0)-R₁₀, wherein R₁₀ is -Ar₃ and the Ar₃ cyclic group is 4-(dimethylaminomethyl)phenyl, 4-(N-morpholinomethyl)phenyl, 4-(N-

25 methylpiperazino)methyl)phenyl, 4-(N-(2methyl)imidazolylmethyl)phenyl, 5-benzimidazolyi, 5benztriazolyl, N-carboethoxy-5-benztriazolyl, Ncarboethoxy-5-benzimidazolyl, or

-C(0)-OR₉, wherein R_9 is -CH₂-Ar₃, and the Ar₃ cyclic group is phenyl, unsubstituted by -Q₁,; and when

 $\rm Y_2$ is $\rm H_2$, $\rm R_3$ is -C(0)-CH $_2$ -T $_1$ -R $_{11}$, T $_1$ is O, and R $_{11}$

- 882 -

is $-C(0)-Ar_4$, wherein the Ar_4 cyclic group is 2,5-dichlorophenyl, then R_5 cannot be:

-C(0)-OR9, wherein ${\rm R}_9$ is -CH2-Ar3 and the ${\rm Ar}_3$ cyclic group is phenyl.

- 5 69. The compound according to claim 68, wherein R_{21} is ${\mbox{-CH}}_3.$
 - 70. The compound according to claim 68, wherein R_{5} is $-C\left(0\right)-C\left(0\right)-OR_{10}$.
- 71. The compound according to claim 68, wherein R_5 is -C(0)-C(0)-OR₁₀ and R_{21} is -CH₃.
 - 72. The compound according to any one of claims 66, 67, 70 and 71, wherein R_3 is -C(0)-H.
 - 73. The compound according to any one of claims 65, 68 and 69, wherein R_3 is -C(0)-H.
- 74. The compound according to claim 68, wherein:

 R_3 is -C(0)-H, and

 R_5 is $-C(0)-R_{10}$, wherein:

 R_{10} is Ar_3 , wherein the Ar_3 cyclic group is phenyl optionally being singly or multiply substituted by:

-F,

-C1,

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 $-N(H)-R_5$, wherein $-R_5$ is -H or $-C(0)-R_{10}$, wherein R_{10} is a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein Ar_2 is

- 883 -

phenyl,

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 $^{-N}(R_9)$ (R_{10}), wherein R_9 and R_{10} are independently a $^{-C}_{1-4}$ straight or branched alkyl group, or $^{-O-R_5}$, wherein R_9 is H or a $^{-C}_{1-4}$ straight or branched alkyl group.

- 75. The compound according to claim 74, wherein Ar_3 is phenyl being optionally singly or multiply substituted at the 3- or 5-position by -Cl or at the 4-position by -NH-R₅, -N(R₃)(R_{10}), or -O-R₅.
- 76. The compound according to claim 68, wherein:

$$R_3$$
 is $-C(0)-H;$

 R_5 is -C(0)- R_{10} , wherein R_{10} is Ar_3 and the Ar_3 cyclic group is selected from the group consisting of is indolyl, benzimidazolyl, thienyl, and benzo(b)thiophenyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.

 $\,$ 77. The compound according to claim 68, wherein:

 $\rm R_5$ is -C(0)-R₁₀, wherein $\rm R_{10}$ is Ar₃ and the Ar₃ cyclic group is selected from quinolyl and isoquinolyl, and said cyclic group optionally being singly or multiply substituted by -Q₁.

78. The compound according to claim 68, wherein:

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 R_3 is -C(0)-H; and

 $\rm R_{5}$ is -C(O)-R $_{10},$ wherein $\rm R_{10}$ is Ar $_{3}$ and the Ar $_{3}$ cyclic group is phenyl, substituted by

O / \ CH₂

\$79.\$ The compound according to claim 68, selected from the group consisting of:

2201 OME COOH COOH

80. A compound represented by the formula:

15 (VI) R₁-N-1

wherein:

5

R₁ 1s:



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(e10)
$$\begin{array}{c} R_{2} \\ R_{5} - N \\ H \end{array} \hspace{0.5cm} , \hspace{0.5cm} \text{or} \hspace{0.5cm} \end{array}$$

C is a ring chosen from the set consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl; the ring optionally being singly or multiply substituted by -01;

10 R₂ is:

(a)
$$(pm)_{R_{S_1}}, \text{ or } (b)$$

m is 1 or 2;

15 each R_5 is independently selected from the group consisting of:

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 $\begin{array}{c} -S\left(O\right)_{2}-R_{9}, \\ -S\left(O\right)_{2}-NH-R_{10}, \\ -C\left(O\right)-CH_{2}-O-R_{9}, \\ -C\left(O\right)C\left(O\right)-R_{10}, \\ \end{array}$ $\begin{array}{c} -R_{9}, \\ -H, \\ -C\left(O\right)C\left(O\right)-OR_{10}, \text{ and } \\ -C\left(O\right)C\left(O\right)-N\left(R_{9}\right)\left(R_{10}\right); \end{array}$

 X_5 is CH or N;

Y2 is H2 or O;

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 $$R_{6}$$ is selected from the group consisting of -H and 15 $$-\text{CH}_{3}$;}$

 $\begin{array}{c} R_8 \text{ is selected from the group consisting of:} \\ -C(O)-R_{10}, \\ -C(O)-CR_{9}, \\ -C(O)-N(H)-R_{10}, \\ \\ 20 & -S(O)_2-R_{9}, \\ -S(O)_2-NH-R_{10}, \\ -C(O)-CH_2-OR_{10}, \\ -C(O)-CH_2-OR_{10}, \\ -C(O)-CH_2N(R_{10})(R_{10}), \\ \\ 25 & -C(O)-CH_2C(O)-CR_{9}, \\ -C(O)-CH_2C(O)-R_{9}, \\ -H, \text{ and } \\ -C(O)-C(O)-CR_{10}; \end{array}$

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

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- 887 -

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

 $\rm R_{13}$ is selected from the group consisting of H, Ar₃, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₉, or -CO₂H;

each R_{51} is independently selected from the group consisting of R_9 , $-C(0)-R_9$, $-C(0)-N(H)-R_9$, or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

each R₂₁ is independently selected from the group consisting of -H or a -C₁₋₆ straight or branched alkyl group;

each Ar_3 is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO_2 , =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by $-Q_1$:

each Q_1 is independently selected from the group

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consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN, =0, -OH, -perfluoro C_{1-3} alkyl, R_5 , -OR₅, -NHR₅, -OR₉, -N(R_9)(R_{10}), - R_9 , -C(O)- R_{10} , and O CH₂,

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

81. The compound according to claim 80, wherein:

15 m is 1;

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1.0

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C is a ring chosen from the set consisting of benzo, pyrido, or thieno the ring optionally being singly or multiply substituted by halogen, $-NH_2$, $-NH-R_5$, $-NH-R_9$, $-OR_{10}$, or $-R_9$, wherein R_9 is a straight or branched C_{1-4} alkyl group, and R_{10} is H or a straight or branched C_{1-4} alkyl group;

Re is H;

 R_{13} is H or a C_{1-4} straight or branched alkyl group optionally substituted with $-Ar_3,\,\,-OH,\,\,-OR_9,\,\,-CO_2H,$ wherein the R_9 is a C_{1-4} branched or straight chain alkyl group; wherein Ar_3 is morpholinyl or phenyl, wherein the phenyl is optionally substituted by $-Q_1;$

 R_{21} is -H or -CH₃;

 R_{51} is a C_{1-6} straight or branched alkyl group

- 889 -

optionally substituted with -Ar₃, wherein Ar₃ is phenyl, optionally substituted by $-O_1$;

each Ar $_3$ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$;

each $\rm Q_1$ is independently selected from the group consisting of -NH2, -Cl, -F, -Br, -OH, -R9, -NH-R5

wherein R_5 is $-C(0)-R_{10}$ or $-S(0)_2-R_9$, $-OR_5$ wherein R_5 is $-C(0)-R_{10}$, $-OR_9$, $-NHR_9$, and

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CH₂,

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

82. The compound according to claim 81, wherein $R_{\rm 1}$ is (w2).

83. The compound according to claim 82,

- 890 -

selected from the group consisting of:

- 84. The compound according to claim 82,
- wherein R₈ is selected from the group consisting of:
 - -C(0)-R10,
 - -C(0)0-Rq,
 - -C(0)-CH2-OR10, and
 - -C(0)-CH2C(0)-R9.
- 10 85. The compound according to claim 84, wherein R_8 is -C(0)-CH₂-OR₁₀ and R_{10} is -H or -CH₃.
 - 86. The compound according to claim 81, wherein ${\bf R}_1$ is (e10) and ${\bf X}_5$ is CH.
- 87. The compound according to claim 81, 15 wherein R_1 is (e10) and $X_{\scriptscriptstyle\rm E}$ is N.
 - 86. The compound according to any one of claims 80-87 wherein $\rm R_5$ is -C(0)-R₁₀ or -C(0)-C(0)-R₁₀.

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89. The compound according to claim 88, wherein $R_{1,0}$ is $\mbox{Ar}_{3}.$

90. The compound according to claim 89, wherein:

 R_5 is -C(O)- R_{10} and R_{10} is Ar_3 , wherein the Ar_3 cyclic group is phenyl optionally being singly or multiply substituted by:

-R₉, wherein R₉ is a C_{1-4} straight or branched alkyl group;

10 -F,

5

15

-Cl,

 $-N(H)-R_5$, wherein $-R_5$ is -H or $-C(0)-R_{10}$, wherein R_{10} is a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein Ar_3 is phenyl,

-N(Rg)(R10), wherein R9 and R10 are independently a -C1-4 straight or branched alkyl group, or

=0-R5, wherein R5 is H or a -C $_{1-4}$ straight or branched alkyl group.

91. The compound according to claim 90, selected from the group consisting of:

- 892 -

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92. The compound according to claim 90, wherein Ar_3 is phenyl being singly or multiply substituted at the 3- or 5-position by -C1 or at the 4-position by -NH-R₅, -N(R₉) (R₁₀), or -O-R₅.

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93. The compound according to claim 92, selected from the group consisting of:

94. The compound according to claim 92, selected from the group consisting of:

- 894 -

95. The compound according to claim 90, wherein Ar_3 is phenyl being singly or multiply substituted at the 3- or 5-position by $\mbox{-Rg},$ wherein \mbox{Rg} is a \mbox{C}_{1-4} straight or branched alkyl group;

- 895 -

and at the 4-position by $-0-R_5$.

96. The compound according to claim 95, selected from the group consisting of:

- 896 -

97. The compound according to claim 95, selected from the group consisting of:

- 897 -

98. The compound according to claim 89,

5 wherein:

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 R_{5} is -C(0)- $R_{10},$ wherein R_{10} is Ar_{3} and the Ar_{3} cyclic group is selected from the group consisting of is indolyl, benzimidazolyl, thienyl, quinolyl, isoquinolyl and benzo(b)thiophenyl, and said cyclic group optionally being singly or multiply substituted by -Q₁.

99. The compound according to claim 98, wherein the Ar_3 cyclic group is isoquinolyl, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.

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100. The compound according to claim 99 selected from the group consisting of:

- 899 -

- 900 -

101. The compound according to claim 99, selected from the group consisting of:

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- 901 -

5 412g

102. The compound according to claim 89, wherein R_5 is -C(0)- R_{10} , wherein R_{10} is Ar_3 and the Ar_3 cyclic group is phenyl, substituted by

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103. The compound according to claim 102, selected from the group consisting of:

- 903 -

- 904 -

104. A compound represented by the formula:

wherein:

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m is 1 or 2;

 $\ensuremath{R_{1}}$ is selected from the group consisting of the following formulae:

C is a ring chosen from the set consisting of benzo, pyrido, thieno, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl, the ring optionally being singly or multiply substituted by -01,;

 R_3 is selected from the group consisting of: -CN, -C(0)-H, -C(0)-CH₂-T₁-R₁₁, -C(0)-CH₂-F', -C=M-O-R₉, and -CO-Ar,;

each $\ensuremath{R_{5}}$ is independently selected from the group consisting of:

- 905 -

$$\begin{array}{c} -C\left(O\right) \, O - R_9, \\ -C\left(O\right) \, -N\left(R_{10}\right) \, \left(R_{10}\right) \\ -S\left(O\right) \, {}_2 \, R_9, \\ -S\left(O\right) \, {}_2 \, -NH - R_{10}, \\ \end{array}$$

$$\begin{array}{c} -C\left(O\right) \, -CH_2 - O - R_9, \\ -C\left(O\right) \, C\left(O\right) - R_{10}, \\ -R_9, \\ -H, \\ -C\left(O\right) \, C\left(O\right) - OR_{10}, \text{ and} \\ -C\left(O\right) \, C\left(O\right) - N\left(R_9\right) \, \left(R_{10}\right); \end{array}$$

each T $_1$ is independently selected from the group consisting of -O-, -S-, -S(O)-, and -S(O) $_2-$;

 $$\rm 15$$ $$\rm R_{6}$$ is selected from the group consisting of -H and -CH $_{\rm 3};$

 $\ensuremath{R_8}$ is selected from the group consisting of:

$$\begin{array}{c} -C(0) - R_{10}, \\ -C(0) - R_{9}, \\ \\ -C(0) - R_{9}, \\ \end{array}$$
 20
$$\begin{array}{c} -C(0) - NH - R_{10}, \\ -S(0) _{2} - R_{9}, \\ -S(0) _{2} - NH - R_{10}, \\ -C(0) - CH_{2} - OR_{10}, \\ -C(0) - CH_{2} - N(R_{10}) (R_{10}), \\ -C(0) - CH_{2} - C(0) - CH_{2} - C(0) - CH_{2} - C(0) - CH_{2} - C(0) - CH_{2} - C(0), \\ -C(0) - CH_{2} - C(0) - CH_{2} - C(0) - CH_{2}, \\ -C(0) - CH_{2} - C(0) - CH_{2} - C(0) - CH_{2}, \\ -C(0) - CH_{2} - C(0) - CH_{2} -$$

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each $\rm R_9$ is independently selected from the group consisting of -Ar $_3$ and a -C $_{1-6}$ straight or branched

-C(0)-C(0)-OR10;

- 906 -

alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

each $\ensuremath{R_{11}}$ is independently selected from the group consisting of:

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 R_{15} is selected from the group consisting of -OH, -OAr₃, -N(H)-OH, and -OC₁₋₆, wherein C_{1-6} is a straight or branched alkyl group optionally substituted with -Ar₃, -CONH₂, -OR₅, -OH, -OR₆, or -CO₂H;

 Ar_2 is independently selected from the following group, in which any ring may optionally be singly or multiply substituted by $-Q_1$ or phenyl, optionally substituted by O_1 :

$$(hh)$$
 , and

- 907 -



wherein each Y is independently selected from the group consisting of O and S;

each Ar₃ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO₂, =N-, and -NH-, -N(R₅)-, and -N(R₉)- said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -O₃:

each Ar $_4$ is a cyclic group independently selected from the set consisting of an aryl group which contains 6, 10, 12, or 14 carbon atoms and between 1 and 3 rings, and a heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -O-, -S-, -SO-, SO $_2$, =N-, -NH-, -N(R $_5$)-, and -N(R $_9$)- said heterocycle group optionally containing one or more double bonds, said heterocycle group cyclic group optionally comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -Q $_1$;

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each Q_1 is independently selected from the group consisting of -NH₂, -CO₂H, -Cl, -F, -Br, -I, -NO₂, -CN,

- 908 -

=C, -OH, -perfluoro
$$C_{1-3}$$
 alkyl, R_5 , -OR $_5$, -NHR $_5$, -OR $_9$, -N(R_9)(R_{10}), - R_9 , -C(O)- R_{10} , and O CH $_2$;

provided that when $-Ar_3$ is substituted with a Q_1 group which comprises one or more additional $-Ar_3$ groups, said additional $-Ar_3$ groups are not substituted with another $-Ar_3$.

105. The compound according to claim 104, wherein:

m is 1:

C is a ring chosen from the set consisting of benzo, pyrido, and thieno, the ring optionally being singly or multiply substituted by halogen, $-{\rm NH}_2$, $-{\rm NH-R}_5$, or $-{\rm NH-R}_9$, $-{\rm OR}_{10}$, or $-{\rm R}_9$, wherein ${\rm R}_9$ is a straight or branched ${\rm C}_{1-4}$ alkyl group, and ${\rm R}_{10}$ is H or a straight or branched ${\rm C}_{1-4}$ alkyl group;

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R6 is H;

 $\rm R_{11}$ is selected from the group consisting of -Ar_4, -(CH_2)_{1=3}-Ar_4, and -C(O)-Ar_4;

Y is O;

- 909 -

each Ar₃ cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, thiazolyl, benzimidazolyl, thienothienyl, thiadiazolyl, benzotriazolyl, benzo[b]thiophenyl, benzofuranyl, and indolyl, and said cyclic group optionally being singly or multiply

each Ar₄ cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, naphthyl, pyridinyl, oxazolyl, pyrimidinyl, or indolyl, and said cyclic group optionally being singly or multiply substituted by -0;

each Q₁ is independently selected from the group consisting of -NH₂, -Cl, -F, -Br, -OH, -R₉, -NH-R₅ wherein R₅ is -C(0)-R₁₀ or -S(0)₂-R₉, -OR₅ wherein R₅ is -C(0)-R₁₀, -NHR₀, and

substituted by -Q1;

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3.0

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

106. The compound according to claim 105, wherein R_8 is selected from the group consisting of:

- 910 -

 $-C(0) -R_{10}$, $-C(0) O -R_{9}$, $-C(0) -CH_2 -OR_{10}$, and $-C(0) -CH_2C(0) -R_{9}$.

107. The compound according to claim 106, wherein R_8 is -C(O)-CH₂-OR₁₀ and R_{10} is -H or -CH₃.

108. The compound according to claim 105, wherein ${\rm R}_3$ is -C(O)-Ar_2,

 $109. \label{eq:compound} 109. \label{eq:compound} The compound according to claim 105, \\ 10 \qquad wherein R_3 is -C(O)CH_2-T_1-R_{11};$

110. The compound according to claim 105, wherein $\rm R_3$ is -C(O)-H.

 $\mbox{111.} \ \mbox{ The compound according to claim 110,} \\ \mbox{wherein R_8 is selected from the group consisting of:}$

-C(O)-R₁₀,

5

15

-C(0)0-Rg,

-C(0) $-CH_2$ $-OR_{10}$, and

-C(O)-CH2C(O)-R9.

112. The compound according to claim 111, 20 selected from the group consisting of:

- 911 -

- 912 -

- 913 -

113. The compound according to claim 111, wherein R_8 is $-C(0)-CH_2-OR_{10}$ and R_{10} is -H or $-CH_3$.

114. The compound according to claim 68,

wherein: m is 1:

5

15

20

T; is 0 or S;

R₂₁ is -H or -CH₃;

10 Aro is (hh):

Y is O:

each Ar3 cyclic group is independently selected from the set consisting of phenyl, naphthyl, thienyl, quinolinyl, isoquinolinyl, pyrazolyl, thiazolyl, isoxazolyl, benzotriazolyl, benzimidazolyl, thienothienyl, imidazolyl, thiadiazolyl, benzo[b]thiophenyl, pyridyl, benzofuranyl, and indolyl and said cyclic group being singly or multiply substituted by $-Q_1$;

each Ar4 cyclic group is independently selected from the set consisting of phenyl, tetrazolyl, pyridinyl, oxazolyl, naphthyl, pyrimidinyl, and thienyl

- 914 -

and said cyclic group being singly or multiply substituted by $-Q_1$;

each Q₁ is independently selected from the group 5 consisting of $-NH_2$, -Cl, -F, -Br, -OH, $-R_9$, $-NH-R_5$ wherein R_5 is $-C(O)-R_{10}$ or $-S(O)_2-R_9$, $-OR_5$ wherein R_5 is $-C(O)-R_{10}$, $-OR_9$, $-NHR_9$, and

CH₂,

10

15

20

25

wherein each R_9 and R_{10} are independently a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$ wherein Ar_3 is phenyl;

provided that when -Ar $_3$ is substituted with a \mathbb{Q}_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$.

- 115. The compound according to claim 114, wherein ${\tt R}_3$ is -C(O)-Ar2,
- . 116. The compound according to claim 114, wherein R_3 is $-C(0)\,CH_2-T_1-R_{11};$
- 117. The compound according to claim 114, wherein R_3 is -C(0)-H.
 - 118. The compound according to any one of claims 104-117, wherein R_5 is -C(0)- R_{10} or -C(0)C(0)- R_{10} .

- 915 -

. 119. The compound according to claim 118, wherein R_{10} is $\mbox{Ar}_3.$

120. The compound according to claim 119, wherein:

 R_5 is -C(O)- R_{10} and R_{10} is Ar_3 , wherein the Ar_3 cyclic group is phenyl optionally being singly or multiply substituted by:

 $-R_9$, wherein R_9 is a C_{1-4} straight or branched alkyl group;

10 -F,

5

15

-Cl,

 $^{-\rm N\,(H)}$ -R₅, wherein -R₅ is -H or -C(0)-R₁₀, wherein R₁₀ is a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein Ar₃ is phenyl,

 $^{-N}\left(R_9\right)\left(R_{10}\right),$ wherein R_9 and R_{10} are independently a $^{-C}_{1-4}$ straight or branched alkyl group, or

 $-\text{O}\text{-R}_5,$ wherein R_5 is H or a $-\text{C}_{1-4}$ straight or branched alkyl group.

20 121. The compound according to claim 120, selected from the group consisting of:

- 916 -

- 917 -

5

122. The compound according to claim 120, wherein Ar_3 is phenyl being singly or multiply substituted at the 3- or 5-position by -Cl or at the 4-position by -NH-R₅, -N(R₉) (R₁₀), or -O-R₅.

123. The compound according to claim 122, selected from the group consisting of:

- 918 -

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124. The compound according to claim 122, 5 selected from the group consisting of:

- 920 -

214m CH₀O H OH

125. The compound according to claim 120, wherein Ar_3 is phenyl being singly or multiply substituted at the 3- or 5-position by $-R_9$, wherein R_9 is a C_{1-4} straight or branched alkyl group; and at the 4-position by $-0-R_5$.

126. The compound according to claim 125, selected from the group consisting of:

10

- 921 -

;

127. The compound according to claim 125, wherein the compound is:

5 214w

10

\$128.\$ The compound according to claim 119, wherein:

 $\rm R_{5}$ is -C(0)-R₁₀, wherein R₁₀ is Ar₃ and the Ar₃ cyclic group is selected from the group consisting of is indolyl, benzimidazolyl, thienyl, quinolyl, isoquinolyl and benzo(b|thiophenyl, and said cyclic group optionally being singly or multiply substituted

- 923 -

by $-Q_1$.

129. The compound according to claim 128, selected from the group consisting of:

- 130. The compound according to claim 128, wherein the Ar $_3$ cyclic group is isoquinoly1, and said cyclic group optionally being singly or multiply substituted by $-Q_1$.
- 10 l31. The compound according to claim 130, wherein the compound is:

- 924 -

 $\mbox{132. The compound according to claim 130,} \label{eq:132}$ wherein the compound is:

133. The compound according to claim 119, wherein R_5 is -C(0)- $R_{10},$ wherein R_{10} is Ar_3 and the Ar_3

- 925 -

cyclic group is phenyl, substituted by

5

15

134. The compound according to claim 133, wherein the compound is:

10 \$135.\$ The compound according to claim 133, wherein the compound is:

136. A pharmaceutical composition, comprising a compound according to any one of claims 1-41 and 57-135 in an amount effective for decreasing IGIF production and a pharmaceutically acceptable carrier.

137. A pharmaceutical composition comprising a compound according to any one of claims 1-41 and 57-135 in an amount effective for decreasing IFN-y

- 926 -

production and a pharmaceutically acceptable carrier.

138. A method for treating or preventing a disease selected from an IGIF mediated disease, an IFN-y mediated disease, an inflammatory disease, an 5 autoimmune disease, an infectious disease, a proliferative disease, a neurodegenerative disease, a necrotic disease, osteoarthritis, acute pancreatitis, chronic pancreatitis, asthma, rheumatoid arthritis. inflammatory bowel disease, Crohn's disease, ulcerative 10 collitis, cerebral ischemia, myocardial ischemia, adult respiratory distress syndrome, infectious hepatitis, sepsis, septic shock, Shigellosis, glomerulonephritis, systemic lupus erythematosus, scleroderma, chronic thyroiditis, Graves' disease, autoimmune gastritis, 15 insulin-dependent diabetes mellitus (Type I), juvenile diabetes, autoimmune hemolytic anemia, autoimmune neutropenia, thrombocytopenia, myasthenia gravis, multiple sclerosis, psoriasis, lichenplanus, graft vs. host disease, acute dermatomyositis, eczema, primary 20 cirrhosis, hepatitis, uveitis, Behcet's disease, acute dermatomyositis, atopic skin disease, pure red cell aplasia, aplastic anemia, amyotrophic lateral sclerosis and nephrotic syndrome comprising the step of administering to said patient a pharmaceutical 25 composition according to claims 136 or 137.

139. The method according to claim 138, wherein the disease is selected from an inflammatory disease, an autoimmune disease, an infectious disease, rheumatoid arthritis, ulcerative collitis, Crohn's disease, hepatitis, adult respiratory distress syndrome, glomerulonephritis, insulin-dependent

3.0

- 927 -

diabetes mellitus (Type I), juvenile diabetes, psoriasis, graft vs. host disease, and hepatitis.

- 140. A process for preparing an N-acylamino compound, comprising the steps of:
- a) mixing a carboxylic acid with an Nalloc-protected amine in the presence of an inert solvent, triphenylphoshine, a nucleophilic scavenger, and tetrakis-triphenyl phosphine palladium(0) at ambient temperature under an inert atmosphere; and
- b) adding to the step a) mixture, HoBT and EDC; and optionally comprising the further step of:

15

25

- c) hydrolyzing the step b) mixture in the presence of a solution comprising an acid and H_2O , wherein the step b) mixture is optionally concentrated.
 - 141. The process according to claim 140, wherein the inert solvent is CH_2Cl_2 , DMF, or a mixture of CH_2Cl_2 and DMF.
- 142. The process according to claim 140, wherein the nucleophilic scavenger is dimedone, morpholine, trimethylsilyl dimethylamine or dimethyl barbituric acid.
 - 143. The process according to claim 142, wherein the nucleophilic scavenger is trimethylsilyl dimethylamine or dimethyl barbituric acid.
 - 144. The process according to claim 142, wherein the inert solvent is CH_2Cl_2 , DMF, or a mixture

- 928 -

of CH2Cl2 and DMF.

- 145. The process according to claim 144, wherein the nucleophilic scavenger is dimethyl barbituric acid.
- 5 146. The process according to claim 145, wherein the solution comprises trifluoroacetic acid in about 1-90% by weight.
 - 147. The process according to claim 146, wherein the solution comprises trifluoroacetic acid in about 20-50% by weight.
 - 148. The process according to claim 145, wherein the solution comprises hydrochloric acid in about 0.1-30% by weight.
- 149. The process according to claim 148, 15 wherein the solution comprises hydrochloric acid in about 5-15% by weight.
 - 150. The process according to any one of claims 140-149, wherein the N-acylamino compound is represented by formula (VIII):

wherein:

10

 $$R_{1}$$ is selected from the group consisting of the following formulae:

- 929 -

5
$$(e12)$$
 R_{21} R_{21}

- 930 -

C is a ring chosen from the set consisting of benzo, pyrido, thiero, pyrrolo, furano, thiazolo, isothiazolo, oxazolo, isoxazolo, pyrimido, imidazolo, cyclopentyl, and cyclohexyl, the ring optionally being singly or multiply substituted by halogen, $-NH_2$, or $-NH-R_0$;

R₂ is:

5

15 m is 1 or 2;

each $R_{\bar{\mathbf{5}}}$ is independently selected from the group consisting of:

- 931 -

```
-C(O)O-Ra,
                      -C(0)-N(R_{10})(R_{10})
                      -S(O)2-Rg,
                      -S(0)2-NH-R10,
                     -C(O)-CH<sub>2</sub>-O-R<sub>9</sub>,
  5
                     -C(0)C(0)-R<sub>10</sub>,
                      -R9.
                     -H,
                     -C(0)C(0)-OR_{10}, and
10
                     -C(0)C(0)-N(R9)(R10);
               X_5 is CH or N;
               Y2 is H2 or O;
               X_7 is -N(R_8) - or -O-;
15
               R_{\rm 6} is selected from the group consisting of -H and
        -CH3;
               R_{\mbox{\scriptsize R}} is selected from the group consisting of:
                     -C(O)-R<sub>10</sub>,
                     -C(0)0-Rg,
20
                     -C(0)-N(H)-R10,
                     -S(0)2-Rg,
                     -S(O)2-NH-R10,
                     -C(O)-CH2-OR10,
25
                     -C(0)C(0)-R10;
                     -C(0) - CH_2N(R_{10})(R_{10}),
                     -C(0)-CH2C(0)-O-R9,
                     -C(O)-CH2C(O)-Rg,
                     -H, and
                    -C(O)-C(O)-OR<sub>10</sub>;
30
```

- 932 -

each R_9 is independently selected from the group consisting of $-Ar_3$ and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, wherein the $-C_{1-6}$ alkyl group is optionally unsaturated;

5 each R_{10} is independently selected from the group consisting of -H, -Ar₃, a -C₃₋₆ cycloalkyl group, and a -C₁₋₆ straight or branched alkyl group optionally substituted with -Ar₃, wherein the -C₁₋₆ alkyl group is optionally unsaturated;

10 R_{13} is selected from the group consisting of H, Ar₃, and a $-C_{1-6}$ straight or branched alkyl group optionally substituted with $-Ar_3$, $-CONH_2$, $-OR_5$, -OH, $-OR_9$, or $-CO_2H$;

15

20

2.5

each R_{51} is independently selected from the group consisting of R_9 , $-C(0)-R_9$, $-C(0)-N(H)-R_9$, or each R_{51} taken together forms a saturated 4-8 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

each R_{21} is independently selected from the group consisting of -H or a $-C_{1-6}$ straight or branched alkyl group;

each ${\rm Ar_3}$ is a cyclic group independently selected from the set consisting of an aryl group which contains ϵ , 10, 12, or 14 carbon atoms and between 1 and 3 rings and an aromatic heterocycle group containing between 5 and 15 ring atoms and between 1 and 3 rings, said heterocyclic group containing at least one heteroatom group selected from -0-, -S-, -S0-, S0₂, =N-, and -NH-, said heterocycle group optionally containing one or more double bonds, said heterocycle group optionally

- 933 -

comprising one or more aromatic rings, and said cyclic group optionally being singly or multiply substituted by -0_1 ;

10 CH₂,

provided that when -Ar $_3$ is substituted with a Q_1 group which comprises one or more additional -Ar $_3$ groups, said additional -Ar $_3$ groups are not substituted with another -Ar $_3$;

151. The process according to any one of claims 140 -149 wherein the N-alloc protected amine is:

Alloc—N OR,

20

25

 R_{51} is independently selected from the group consisting of R_9 , $-C(O)-R_9$, $-C(O)-N(H)-R_9$, or each R_{51} taken together forms a saturated 4-6 member carbocyclic ring or heterocyclic ring containing -O-, -S-, or -NH-;

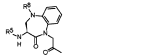
152. The process according to any one of claims 140-149, wherein R_1 is:

- 934 -

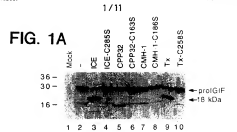
(A-e10)

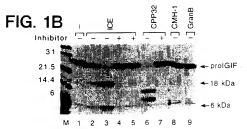
153. The process according to any one of claims 140-149, wherein $\ensuremath{R_{1}}$ is:

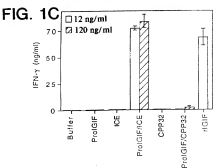
5 (A-w2)

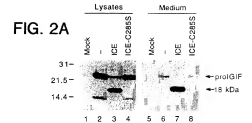


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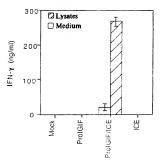


FIG. 2B

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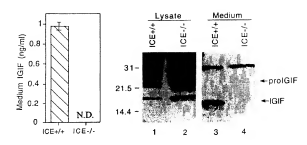


FIG. 3A

FIG. 3B

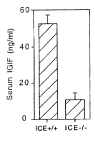


FIG. 3C

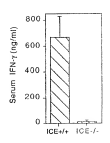
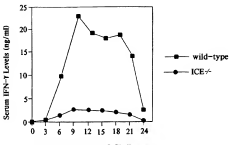


FIG. 3D



Time (hours) after LPS Challenge

FIG. 4

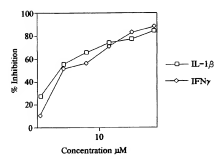
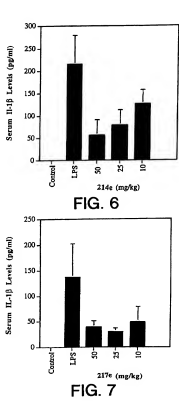


FIG. 5



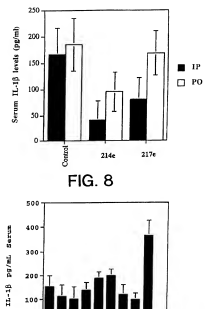


FIG. 9

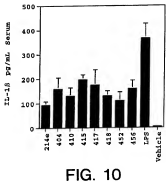
417

404-410-415-

452

LPS

Vehicle



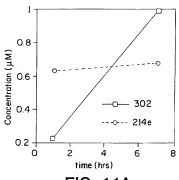


FIG. 11A

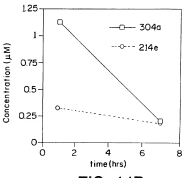


FIG. 11B

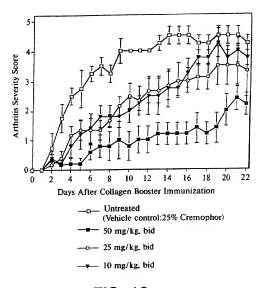
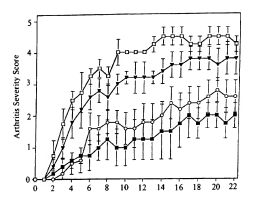


FIG. 12



Days After Collagen Booster Immunization

_____ Untreated
(Vehicle control:25% Cremophor)

---- 50 mg/kg, bid

-o- 25 mg/kg, bid

_____ 10 mg/kg, bid

FIG. 13

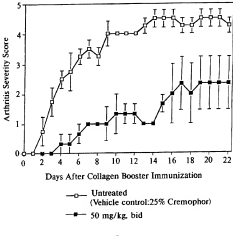


FIG. 14

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(a), and made and a second of the second of		ham, MA 02194 (US). MURCKO, Mark, A.; 520 Marshall
(30) Priority Data: (80,573,641 20 December 1995 (20.12.9) (80,573,642 21 E February 1996 (20.12.9) (80,573,278 12 E Spermher 1996 (20.12.9) (80,013,495 26 November 1996 (26.11.9) (80,761,483 6 December 1996 (06.12.96)	(106) t (6) t (7) t	Street, Holliston, M. A. 01746 (US). MURDOCH, Robert. 89 (Knowlands, Highworth, Wilshine SNG 7ND (GB). NYC. 152 Philip, L.: 18 Praspect Street, Millbury, MA. 01527 (US). ROBIDOUX, Andrea, L., C.; 180 Salem Street, Andower. 153 MA. 01810 (US). SU, Michael, 15 Donna Road, Newton, MA. 02159 (US). WANNAMAKER, M., Woods; 375 Harvard Road, Stow, MA. 01775 (US). WILSON, Keith, P.; 6 Longwood Drive, Hopkinton, MA. 01745 (US). ZELLE, Robert, E.; 67 Boon Road, Stow, MA. 01775 (US).
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(57) Abstract

The present invention relates to novel classes of compounds which are inhibitors of interleukin- 1β converting enzyme. The ICE inhibitors of this invention are characterized by specific structural and physicochemical features. This invention also relates to pharmaceutical compositions comprising these compounds. The compounds and pharmaceutical compositions of this invention are particularly well suited for inhibiting ICE activity and consequently, may be advantageously used as agents against IL-1-, apoptosis-, IGIF-, and IFN-y- mediated diseases, inflammatory diseases, autoimmune diseases, destructive bone disorders, proliferative disorders, infectious diseases, degenerative diseases, and necrotic diseases. This invention also relates to methods for inhibiting ICE activity, for treating interleukin-1-, apoptosis-, IGIF- and IFN-y-mediated diseases and decreasing IGIF and IFN-y production using the compounds and compositions of this invention. This invention also relates to methods for preparing N-acylamino compounds.

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Intern. al Application No PCT/US 96/20843

Relevant to claim No.

A. CLASSIFICATION OF SUBJECT MATTER
1PC 6 C07K5/023 C07D487/04 C07D498/04 A61K38/04 A61K31/55 //(C07D487/04,243:00,237:00)

According to International Patent Classification (IPC) or to both national classification and IPC

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Minimum documentation searched (classification system followed by classification symbols) IPC 6 CO7K CO7D

Category * Citation of document, with indication, where appropriate, of the relevant passages

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

CONSIDERED TO	

x	EP 0 135 349 A (TAKEDA CHEMICAL INDUSTRIES LTD) 27 March 1985 see claims 1-4,41-43; examples 24,25	1,42-56, 136-139
	-/	
	-	

Patent family members are listed in annex.

* Special categories of cited documents:

- *A' document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier document but published on or after the international filing date
- 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
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- "I" later document published after the international filing date or priority date and not in conflict with the application bu-cited to understand the principle or theory underlying the invention. "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "vivare as instance kep writin the document is taken alone for document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled in the art.
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Date of the actual completion of the international search Date of mailing of the international search report 0 4, 09, 97

28 August 1997

Authorized officer

Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV R15wn/x Tel. (+31-70) 340-2000, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016

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Intern. at Application No PCT/US 96/20843

	auon) DOCUMENTS CONSIDERED TO BE RELEVANT	Relevant to claim No.
ategory *	Citation of document, with indication, where appropriate, of the relevant passages	
(WO 95 33751 A (SANOFI WINTHROP INC) 14 December 1995 see the whole document	4-8, 16-38, 40-57, 59-62, 64-81, 86-90, 92,94, 95, 97-99, 101-103, 114-120, 122
		127,128, 130,132, 133, 135-139
,х	WO 95 35308 A (VERTEX PHARMA) 28 December 1995 see the whole document	1-8, 14-139
A	EP 0 644 198 A (STERLING WINTHROP INC) 22 March 1995 see the whole document	1-8, 14-139
A	WO 95 26958 A (SANOFI WINTHROP INC) 12 October 1995 cited in the application see the whole document	1-8, 14-139
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PCT/US 96/20843

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This In	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: 55,56,138,139 because they relate to subject matter not required to be searched by this Authority, namely. Remark: Although claim(s) 55,56,138,139 1s(are) directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged
2.	effects of the compound/composition. Claims Nos: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
	cernacional Searching Authoricy found multiple inventions in this international application, as follows: e next page
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all rearchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Noz.:
	Subject 1: Claims 1-13,14,15,39,58,63,82-85,91,93,96,100,104-113,121,123,126, 129,131,134(complete);4-8,42-57,62,80,81,88-90,92,95,98,99,102,118-120,122,125, 128,130,133,136-139(partially). See next page
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/210
Subject 2: Claims 16-38,40,41,59-61,64-79,86,87,94,97,101,103,114-117,124,127, 132,135(complete); 4-8,42-57,62,80,81,88-90,92,95,98,99,102,118-120,122,125, 128,130,133,136-139(partially

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